(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 20 September 2001 (20.09.2001)

(10) International Publication Number WO 01/68864 A1

(51) International Patent Classification7: C12N 15/31, 15/67 // (C12N 15/31, C12R 1:69)

(21) International Application Number: PCT/DK01/00169

(22) International Filing Date: 14 March 2001 (14.03.2001)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data: PA 2000 00406 14 March 2000 (14.03.2000) DK

(71) Applicant: NOVOZYMES A/S [DK/DK]; Krogshøjvej 36, DK-2880 Bagsværd (DK).

(72) Inventors: HJORT, Carsten, M.; Råbroparken 36, DK-2765 Smørum (DK). HONDEL, C., MJJ van den; Waterlelie 124, NL-2804 PZ Gouda (NL). PUNT. P., J.; Boekdukkersgilde, NL-3994 XT Houten (NL). SCHUREN, F., H., J.; Bachlaan 34, NL-3906 ZK Veenendaal (NL). CHRISTENSEN, Tove; Nøddevænget 3, DK-2800 Lyngby (DK).

(81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW.

(84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published:

- with international search report
- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments
- with (an) indication(s) in relation to deposited biological material furnished under Rule 13bis separately from the description

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.





(54) Title: FUNGAL TRANSCRIPTIONAL ACTIVATOR USEFUL IN METHODS FOR PRODUCING POLYPEPTIDES

(57) Abstract: The present invention relates to isolated nucleic acid sequences encoding polypeptides having transcriptional activation activity and to the polypeptides. The invention also relates to nucleic acid constructs, vectors and host cells comprising the nucleic acid sequences. The invention further relates to host cells useful for the production of polypeptides in which the production or function of the transcriptional activator has been altered, as well as to methods for producing the polypeptides.

1

FUNGAL TRANSCRIPTIONAL ACTIVATOR USEFUL IN METHODS FOR PRODUCING POLYPEPTIDES

Background of the Invention

5

Field of the Invention

The present invention relates to isolated nucleic acid sequences encoding polypeptides having transcriptional activation activity and to the polypeptides. The invention also relates to nucleic acid constructs, vectors and host cells comprising the nucleic acid sequences. The invention further relates to host cells useful for the production of polypeptides in which the production or function of the transcriptional activator has been altered, as well as to methods for producing the polypeptides.

Description of the Related Art

The use of recombinant host cells in the expression of heterologous proteins has in recent years greatly simplified the production of large quantities of commercially valuable proteins which otherwise are obtainable only by purification from their native sources. Currently, there is a varied selection of expression systems from which to choose for the production of any given protein, including eubacterial and eukaryotic hosts.

The selection of an appropriate expression system often depends not only on the ability of the host cell to produce adequate yields of the protein in an active state, but, to a large extent, may also be governed by the intended end use of the protein.

One problem frequently encountered is the high level of proteolytic enzymes produced by a given host cell or present in the culture medium. One suggestion has been to provide host organisms deprived of the ability to produce specific proteolytic compounds. For example, WO 90/00192 (Genencor,

2

Inc.) describes filamentous fungal hosts incapable of secreting enzymatically active aspartic proteinase. EP 574 347 (Ciba Geigy AG) describes Aspergillus hosts defective in a serine protease of the subtilisin-type. WO 98/12300 (Novo Nordisk A/S) describes hosts defective in a metalloprotease and an alkaline protease. WO 97/12045 (Genencor, Inc.) describes yeast and bacterial host systems, which are rendered protease deficient resulting from a disruption of a promoter sequence involved in the regulation of a protease gene.

Mattern, I.E., et al., (1992. Mol Gen Genet 234:332-336) describe a mutant strain of Aspergillus niger, which was shown to have only 1 to 2% of the extracellular protease activity of the parent strain, apparently due to a deficiency of at least two proteases, aspergillopepsin A and aspergillopepsin B. It was suggested that the protease deficient phenotype could result from a regulatory mutation affecting the expression of the genes coding for both proteases.

The initiation of eukaryotic transcription at a specific promoter orset of promoters requires а eukaryotic 20 transcriptional activator which is a polypeptide, but which is not itelf part of RNA polymerase. Many transcriptional activators bind to a specific site on the promoter to form a functional promoter necessary for the initiation transcription of the polypeptide encoding sequence. However, a 25 transcriptional activator may also be incorporated into an initiation complex only in the presence of other polypeptides. Polypeptides with transcriptional activation activity have been described in fungi, and a list of such polypeptides has been published (Dhawale, S.S., and Lane, A.C. 1993. Nucleic Acid 30 Research 21:5537-5546).

Solution proposed by the invention:

It is an object of the present invention to provide improved methods for increasing production of polypeptides in host cells

WO 01/68864

3

in which the activity of a transcriptional activator involved in the regulation of protease production has been modified.

Summary of the Invention

10

30

A first aspect of the present invention relates to an isolated nucleic acid sequence encoding a transcriptional activator selected from the group consisting of:

- (a) a nucleic acid sequence having at least 70% identity with the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48;
- (b) a nucleic acid sequence encoding a polypeptide having an amino acid sequence which has at least 50% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49;
- a nucleic acid sequence which hybridizes under low . 15 stringency conditions with (i) the nucleic acid sequence SEQ ID NO:1 or SEQ ID NO: 48, or (ii) complementary strand, wherein the low stringency conditions are defined by prehybridization hybridization at 42°C in 5x SSPE, 0.3% SDS, 200 microg/ml 20 sheared and denatured salmon sperm DNA, and formamide, and wash conditions are defined by 50°C for 30 minutes in 2X SSC, 0.2% SDS;
 - (d) an allelic variant of (a), (b), or (c);
- 25 (e) a subsequence of (a), (b), (c), or (d), wherein the subsequence encodes a polypeptide fragment which has transcriptional activation activity; and
 - (f) a subsequence of (a), (b) (c), or (d), wherein the subsequence encodes a polypeptide with the amino acid sequence of SEQ ID NO:3.

The nucleic acid sequence shown in SEQ ID NO: 1 is the Aspergillus niger prtT gene encoding the transcriptional factor

WO 01/68864

4

shown in SEQ ID NO: 2 as described further below and in the Examples.

The nucleic acid sequence shown in SEQ ID NO: 48 is the Aspergillus oryzae IFO4177 prtT gene encoding the 5 transcriptional factor shown in SEQ ID NO: 49. The A. oryzae prtT gene has a coding region starting in position 795 and ending at position 2931. The prtT gene has 4 introns in positions 1028-1135, 1538-1591, 2018-2066, and 2297-2347, respectively. This is described further below and in the 10 Examples.

In another aspect, the invention also relates to nucleic acid constructs, vectors and host cells comprising the nucleic acid sequences, and to the polypeptides encoded by the nucleic acid sequences. The invention further relates to host cells useful for the production of a polypeptide, in which the production or function of the transcriptional activator has been altered, as well as to methods for producing the polypeptide.

20 Brief Description of the Figures

Figure 1 shows a restriction map of the plasmid pPAP, the construction of which is described in Example 1.

Figure 2 shows a restriction map of the plasmid pAopyrGcosArp1, the construction of which is described in 25 Example 1.

Figure 3 shows a restriction map of the plasmid pEES1, the construction of which is described in Example 1.

Figure 4 shows a restriction map of the plasmid pDprt, the construction of which is described in Example 3.

Figure 5 shows a restriction map of the plasmid pGPprt, the construction of which is described in Example 4.

Figure 6 shows the sequence of the insert in the two plasmids containing the PCR fragment of the $A.\ oryzae\ prtT\ Zn^{2+}$ -

5 .

finger. ICA217 is the sequence from one of the plasmids and ICA218 is the sequence from the other.

Figure 7 shows plasmid pDV8 a pSP65 (Promega™) based plasmid containing the HSV-tk gene on a 1.2 kb BglII/BamHI 5 fragment inserted between a 1.0 kb XhoI/BglII fragment of the A. nidulans gpd promoter and a 0.8 kb BamHI/HindIII fragment containing the A. nidulans trpC transcriptional terminator.

Figure 8 shows the construction of pJaL554 described in Example 8.

10

Detailed Description of the Invention

Nucleic Acid Sequences Encoding Transcriptional Activators

A first aspect of the present invention relates to an isolated nucleic acid sequence encoding a transcriptional activator selected from the group consisting of:

- (a) a nucleic acid sequence having at least 70% identity with the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48;
- 20 (b) a nucleic acid sequence encoding a polypeptide having an amino acid sequence which has at least 50% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49;
- a nucleic acid sequence which hybridizes under low 25 stringency conditions with (i) the nucleic acid sequence SEQ ID NO:1 or SEQ ID NO: 48, or (ii) complementary strand, wherein the low stringency conditions are defined prehybridization by hybridization at 42°C in 5x SSPE, 0.3% SDS, 200 micro 30 g/ml sheared and denatured salmon sperm DNA, and 25% formamide, and wash conditions are defined by 50°C for 30minutes in 2X SSC, 0.2% SDS;
 - (d) an allelic variant of (a), (b), or (c);

6

(e) a subsequence of (a), (b), (c), or (d), wherein the subsequence encodes a polypeptide fragment which has transcriptional activation activity; and

(f) a subsequence of (a), (b), (c), or (d), wherein the subsequence encodes a polypeptide with the amino acid sequence of SEQ ID NO:3.

5

The term "transcriptional activator" as used herein refers to a polypeptide which has the capability to activate a specific promoter or set of promoters necessary for the initiation of transcription of the polypeptide encoding sequence to which it is linked.

The term "isolated nucleic acid sequence" as used herein refers to a nucleic acid sequence which is essentially free of other nucleic acid sequences, e.g., at least about 20% pure, 15 preferably at least about 40% pure, more preferably at least about 60% pure, even more preferably at least about 80% pure, most preferably at least about 90% pure as determined by agarose electrophoresis. For example, an isolated nucleic acid sequence can be obtained by standard cloning procedures used in 20 genetic engineering to relocate the nucleic acid sequence from its natural location to a different site where it will be reproduced. The cloning procedures may involve excision and isolation of a desired nucleic acid fragment comprising the nucleic acid sequence encoding the polypeptide, insertion of 25 the fragment into a vector molecule, and incorporation of the recombinant vector into a host cell where multiple copies or clones of the nucleic acid sequence will be replicated. nucleic acid sequence may be of genomic, cDNA, RNA, semisynthetic, synthetic origin, or any combinations thereof.

In a preferred embodiment, the nucleic acid sequence has a degree of identity to the nucleic acid sequence set forth in SEQ ID NO:1 or SEQ ID NO: 48 of at least about 70%, preferably at least about 80%, more preferably at least about 85%, even more preferably at least about 90%, most preferably at least

7

about 95%, even most preferably at least about 97%, and even more preferred at least 99% identity, which encodes an active polypeptide. For purposes of the present invention, the degree of identity between two nucleic acid sequences is determined by the Clustal method (Higgins, 1989, CABIOS 5:151-153) with an identity table, a gap penalty of 10, and a gap length penalty of 10.

In an even more preferred embodiment, the nucleic acid sequence encoding a transcriptional activator has a nucleic acid sequence as set forth in SEQ ID NO:1 or SEQ ID NO: 48.

Modification of a nucleic acid sequence encoding a polypeptide of the present invention may be necessary for the synthesis of polypeptides substantially similar to the polypep-The term "substantially similar" to the polypeptide 15 refers to non-naturally occurring forms of the polypeptide. These polypeptides may differ in some engineered way from the polypeptide isolated from its native source. For example, it may be of interest to synthesize variants of the polypeptide where the variants differ in specific activity, binding 20 specificity and/or affinity, or the like using, e.g., sitedirected mutagenesis. The analogous sequence may be constructed on the basis of the nucleic acid sequence presented as the polypeptide encoding part of SEQ ID NO: 1 or SEQ ID NO: 48, e.g., a subsequence thereof, and/or by introduction of 25 nucleotide substitutions which do not give rise to another amino acid sequence of the polypeptide encoded by the nucleic acid sequence, but which corresponds to the codon usage of the host organism intended for production of the enzyme, or by introduction of nucleotide substitutions which may give rise to 30 a different amino acid sequence. For a general description of nucleotide substitution, see, e.g., Ford et al., 1991, Protein Expression and Purification 2: 95-107.

In another preferred embodiment, the present invention relates to isolated nucleic acid sequences encoding

8

polypeptides having an amino acid sequence which has a degree of identity to the amino acid sequence set forth in SEQ ID NO:2 or SEQ ID NO: 49 of at least about 50%, preferably at least about 60%, preferably at least about 70%, more preferably at least about 80%, even more preferably at least about 90%, most preferably at least about 95%, even most preferably at least about 97%, and even more preferred at least 99%, which qualitatively retain the transcriptional activation activity of the polypeptides (hereinafter "homologous polypeptides").

- In a preferred embodiment, the homologous polypeptides have an amino acid sequence which differs by five amino acids, preferably by four amino acids, more preferably by three amino acids, even more preferably by two amino acids, and most preferably by one amino acid from the amino acid sequence set forth in SEQ ID NO:2 or SEQ ID NO: 49. For purposes of the present invention, the degree of identity between two amino acid sequences is determined by the Clustal method (Higgins, 1989, supra) with an identity table, a gap penalty of 10, and a gap length penalty of 10.
- 20 Hybridization indicates that by methods of standard Southern blotting procedures, the nucleic acid sequence hybridizes to an oligonucleotide probe corresponding to the polypeptide encoding part of the nucleic acid sequence shown in SEQ ID NO:1 , under low to high stringency conditions (i.e., prehybridization and 25 hybridization at 42°C in 5x SSPE, 0.3% SDS, 200 $\mu g/ml$ sheared and denatured salmon sperm DNA, and either 25, 35 or 50% formamide for low, medium and high stringencies, respectively). In order to identify a clone or DNA which is homologous with SEQ ID NO:1 or SEQ ID NO: 48, the hybridization reaction is 30 washed three times for 30 minutes each using 2X SSC, 0.2% SDS preferably at least 50°C, more preferably at least 55°C, more preferably at least 60°C, more preferably at least 65°C, even more preferably at least 70°C, and most preferably at least 75°C.

9

The nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48, or a subsequence thereof, as well as the amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 49, or a partial sequence thereof, or the amino acid sequence of SEQ ID NO:3, may be used to 5 design an oligonucleotide probe to identify and isolate or clone a homologous gene of any genus or species according to methods well known in the art.

In particular, such probes can be used for hybridization with the genomic or cDNA of the genus or species of interest, following standard Southern blotting procedures, in order to identify and isolate the corresponding gene therein. Such probes can be considerably shorter than the entire sequence, but should be at least 15, preferably at least 25, and more preferably at least 40 nucleotides in length. Longer probes can also be used. Both DNA and RNA probes can be used. The probes are typically labeled for detecting the corresponding gene (for example, with 32P, 3H, 35S, biotin, or avidin). For example, molecules to which a 32P-, 3H- or 35S-labelled oligonucleotide probe hybridizes may be detected by use of X-20 ray film.

Thus, a genomic, cDNA or combinatorial chemical library prepared from such other organisms may be screened for DNA which hybridizes with the probes described above and which encodes a polypeptide with transcriptional activation activity.

25 Genomic or other DNA from such other organisms may be separated by agarose or polyacrylamide gel electrophoresis, or other separation techniques. DNA from the libraries or the separated DNA may be transferred to and immobilized on nitrocellulose or other suitable carrier material. A clone or DNA which is homologous to SEQ ID NO:1 or SEQ ID NO: 48 may then be identified following standard Southern blotting procedures.

An allelic variant denotes any of two or more alternative forms of a gene occupying the same chomosomal locus. Allelic variation arises naturally through mutation, and may result in

10

phenotypic polymorphism within populations. Gene mutations can be silent (i.e., no change in the encoded polypeptide) or may encode polypeptides having altered amino acid sequences.

The term "allelic variant of a polypeptide" is a polypeptide 5 encoded by an allelic variant of a gene. In a preferred embodiment, the nucleic acid sequence encoding transcriptional activator of the present invention is an allelic variant of a nucleic acid sequence selected from the group consisting of nucleic acid sequences: (a) having at least 10 70% identity with the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48, (b) encoding a polypeptide having an amino acid sequence which has at least 50% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49, (c) which hybridizes under low stringency conditions with the nucleic acid sequence 15 of SEQ ID NO:1 or SEQ ID NO: 48, or its complementary strand, and (d) encoding a polypeptide having the amino acid sequence of SEQ ID NO:3.

present invention also The encompasses nucleic acid sequences which differ from SEQ ID NO:1 or SEQ ID NO: 48 by 20 virtue of the degeneracy of the genetic code. The present invention also relates to subsequences of SEQ ID NO:1 or SEQ ID NO: 49, wherein a subsequence of SEQ ID NO:1 is a nucleic acid sequence encompassed by SEQ ID NO:1 or SEQ ID NO: 48 except that one or more nucleotides from the 5' and/or 3' end have 25 been deleted. Preferably, a subsequence of SEQ ID NO:1 or SEQ ID NO: 48 encodes a polypeptide fragment which transcriptional activation activity. In a more preferred embodiment, a subsequence of SEQ ID NO:1 or SEQ ID NO: 48 contains at least a nucleic acid sequence encoding the 30 polypeptide sequence shown in SEQ ID NO:3.

The techniques used to isolate or clone a nucleic acid sequence encoding a polypeptide are known in the art and include isolation from genomic DNA, preparation from cDNA, or a combination thereof. The cloning of the nucleic acid sequences

11

of the present invention from such genomic DNA can be effected, e.g., by using methods based on polymerase chain reaction (PCR) or antibody screening of expression libraries to detect cloned DNA fragments with shared structural features. (See, e.g., 5 Innis et al., 1990, PCR: A Guide to Methods and Application, Academic Press, New York.) Other nucleic acid amplification procedures such as ligase chain reaction (LCR), ligated activated transcription (LAT) and nucleic acid sequence-based amplification (NASBA) may be used. The nucleic acid sequence 10 may be cloned from a microorganism, or another or related organism and thus, for example, may be an allelic or species variant of the polypeptide encoding region of the nucleic acid sequence.

The transcriptional activators encoded by nucleic acid
sequences which hybridize with an oligonucleotide probe which
hybridizes with the nucleic acid sequence of SEQ ID NO:1 or SEQ
ID NO: 48, its complementary strand, or allelic variants and
subsequences of SEQ ID NO:1 or SEQ ID NO: 48, or allelic
variants and fragments of the transcriptional activators may be
obtained from microorganisms of any genus.

In a preferred embodiment, the transcriptional activators may be obtained from a fungal cell. "Fungi" as used herein includes the phyla Ascomycota, Basidiomycota, Chytridiomycota, and Zygomycota (as defined by Hawksworth, et al., in Ainsworth 25 and Bisby's Dictionary of The Fungi, 8th edition, 1995, CAB International, University Press, Cambridge, UK) as well as the Oomycota (as cited in Hawksworth et al., 1995, supra, page 171) and all mitosporic fungi (Hawksworth et al., 1995, supra).

In preferred embodiment, the fungal source is a filamentous fungal strain. "Filamentous fungi" include all filamentous forms of the subdivision Eumycota and Oomycota (as defined by Hawksworth et al., 1995, supra). The filamentous fungi are characterized by a mycelia wall composed of chitin, cellulose, glucan, chitosan, mannan, and other complex polysaccharides.

12

Vegetative growth is by hyphal elongation and carbon catabolism is obligately aerobic. Filamentous fungal strains include, but are not limited to, strains of Acremonium, Aspergillus, Aureobasidium, Cryptococcus, Filibasidium, Fusarium, Humicola, Magnaporthe, Mucor, Myceliophthora, Neocallimastix, Neurospora, Paecilomyces, Penicillium, Piromyces, Schizophyllum, Talaromyces, Thermoascus, Thielavia, Tolypocladium, and Trichoderma.

In a more preferred embodiment, the nucleic acid sequence encoding a transcriptional activator of the present invention is obtained from a strain of Aspergillus, such as A. awamori or A. nidulans. Preferably, the nucleic acid sequence is obtained from a strain of A. niger or A. oryzae. Even more preferably, the nucleic acid sequence is obtained from an isolate of a strain of A. niger, DSM 12298; e.g., the nucleic acid sequence set forth in SEQ ID NO:1, or from A. oryzae IFO 4177, i.e., the nucleic acid sequence set forth in SEQ ID NO: 48.

In another more preferred embodiment, the nucleic acid sequence encoding a transcriptional activator of the present invention is obtained from a strain of Fusarium, such as F. oxysporum. Preferably, the strain is a strain of F. venenatum (Nirenberg sp. nov.).

In another preferred embodiment, the nucleic acid sequence encoding a transcriptional activator of the present invention is obtained from a yeast strain, such as a Candida, Kluyveromyces, Schizosaccharomyces, or Yarrowia strain. Preferably, the strain is a strain of Hansenula, Pichia, or Saccharomyces.

It will be understood that for the aforementioned species, the invention encompasses both the perfect and imperfect states, and other taxonomic equivalents, e.g., anamorphs, regardless of the species name by which they are known. Those skilled in the art will readily recognize the identity of appropriate equivalents. For example, the polypeptides may be

13

obtained from microorganisms, which are taxonomic equivalents of Aspergillus as defined by Raper, K.D. and Fennel, D.I. (1965. The Genus Aspergillus, The Wilkins Company, Baltimore MD). regardless of the species name by which they are known. 5 Aspergilli are mitosporic fungi characterized by an aspergillum comprised of a conidiospore stipe with no known teleomorphic states terminating in a vesicle, which in turn bears one or two layers of synchronously formed specialized cells, variously referred to as sterigmata or phialides, and asexually formed 10 spores referred to as conidia. Known teleomorphs of Aspergillus include Eurotium, Neosartorya, and Emericella. Strains of Aspergillus and teleomorphs thereof are readily accessible to the public in a number of culture collections, such as the American Type Culture Collection (ATCC), Deutsche 15 Sammlung von Mikroorganismen und Zellkulturen GmbH (DSM), Centraalbureau Voor Schimmelcultures (CBS), and Agricultural Research Service Patent Culture Collection, Northern Regional Research Center (NRRL).

Furthermore, such transcriptional activators may be 20 identified and obtained from other sources including microorganisms isolated from nature (e.g., soil, composts, water, etc.) using the above-mentioned probes. Techniques for isolating microorganisms from natural habitats are well known in the art. The nucleic acid sequence may then be derived by 25 similarly screening a genomic or cDNA library of another microorganism. Once a nucleic acid sequence encoding a transcriptional activator has been detected with the probe(s), the sequence may be isolated or cloned by utilizing techniques which are known to those of ordinary skill in the art (see, 30 e.g., J. Sambrook, E.F. Fritsch, and T. Maniatus, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, New York).

In another preferred embodiment, the isolated nucleic acid sequence encodes a polypeptide comprising the amino acid

14

sequence of SEQ ID NO: 2 or SEQ ID NO: 49, or a fragment thereof, which has transcriptional activation activity.

In another preferred embodiment, the isolated nucleic acid sequence encodes a polypeptide comprising the amino acid sequence of SEQ ID NO: 3.

The present invention also relates to isolated nucleic acid sequences encoding a transcriptional activator of the present invention, which, e.g., using methods of standard Southern blotting procedures described above (cf., Sambrook, et al., 1989, supra), hybridize under low stringency conditions, more preferably medium stringency conditions, and most preferably high stringency conditions, with an oligonucleotide probe which hybridizes under the same conditions with the nucleic acid sequence set forth in SEQ ID NO:1 or SEQ ID NO: 48 or its complementary strand, or allelic variants and subsequences of SEQ ID NO:1 or SEQ ID NO: 48 which encode polypeptide fragments which are transcriptional activators in fungi.

In another more preferred embodiment, the nucleic acid sequence is the nucleic acid sequence encoding a polypeptide, which has DNA binding activity contained in the plasmid pEES which is contained in Escherichia coli DSM 12294.

Nucleic Acid Constructs

Another aspect of the present invention relates to nucleic acid constructs comprising a nucleic acid sequence encoding a transcriptional activator of the present invention operably linked to one or more control sequences, which direct the production of the transcriptional activator in a suitable expression host. In a preferred embodiment, the nucleic acid sequence encodes a polypeptide, which is contained in the plasmid pEES harboured in Escherichia coli DSM 12294, or the nucleic acid sequence shown in SEQ ID NO: 48 encoding the polypeptide shown in SEQ ID NO: 49.

15

Expression will be understood to include any step involved in the production of the polypeptide including, but not limited to, transcription, post-transcriptional modification, translation, post-translational modification, and secretion.

5 Manipulation of the nucleic acid sequence encoding a polypeptide prior to its insertion into a vector may be desirable or necessary depending on the expression vector. The techniques for modifying nucleic acid sequences utilizing cloning methods are well known in the art.

"Nucleic acid construct" is defined herein as a nucleic acid 10 molecule, either single- or double-stranded, which is isolated from a naturally occurring gene or which has been modified to contain segments of nucleic acid which are combined and juxtaposed in a manner which would not otherwise exist in 15 nature. The term nucleic acid construct is synonymous with the term expression cassette when the nucleic acid construct contains all the control sequences required for expression of a coding sequence. The term "coding sequence" as defined herein is a sequence, which is transcribed into mRNA and translated 20 into a transcriptional activator of the present invention. boundaries of the coding sequence are generally determined by the ATG start codon at the 5' end of the mRNA and a transcription terminator sequence located just downstream of the open reading frame at the 3' end of the mRNA. A coding 25 sequence can include, but is not limited to, DNA, cDNA, and recombinant nucleic acid sequences.

The term "control sequences" is defined herein to include all components, which are necessary or advantageous for the expression of a polypeptide. Each control sequence may be native or foreign to the nucleic acid sequence encoding the polypeptide. Such control sequences include, but are not limited to, a leader, a polyadenylation sequence, a propeptide sequence, a promoter, a signal sequence, and a transcription terminator. At a minimum, the control sequences include a

16

promoter, and transcriptional and translational stop signals. The control sequences may be provided with linkers for the purpose of introducing specific restriction sites facilitating ligation of the control sequences with the coding region of the nucleic acid sequence encoding a polypeptide. The term "operably linked" is defined herein as a configuration in which a control sequence is appropriately placed at a position relative to the coding sequence of the DNA sequence such that the control sequence directs the production of a polypeptide.

The control sequence may be an appropriate promoter sequence, a nucleic acid sequence, which is recognized by a host cell for expression of the nucleic acid sequence. The promoter sequence contains transcriptional control sequences, which mediate the expression of the polypeptide. The promoter may be any nucleic acid sequence, which shows transcriptional activity in the cell including mutant, truncated, and hybrid promoters, and may be obtained from genes encoding extracellular or intracellular polypeptides either homologous or heterologous to the cell.

The control sequence may also be a suitable transcription terminator sequence, a sequence recognized by a filamentous fungal cell to terminate transcription. The terminator sequence is operably linked to the 3' terminus of the nucleic acid sequence encoding the polypeptide. Any terminator, which is functional in the cell, may be used in the present invention.

Preferred terminators for filamentous fungal cells are obtained from the genes encoding Aspergillus oryzae TAKA amylase, Aspergillus niger glucoamylase, Aspergillus nidulans anthranilate synthase, Aspergillus niger alpha-glucosidase, and Fusarium oxysporum trypsin-like protease.

The control sequence may also be a suitable leader sequence, a nontranslated region of a mRNA which is important for translation by the filamentous fungal cell. The leader

17

sequence is operably linked to the 5' terminus of the nucleic acid sequence encoding the polypeptide. Any leader sequence, which is functional in the cell, may be used in the present invention.

Preferred leaders for filamentous fungal cells are obtained from the genes encoding Aspergillus oryzae TAKA amylase and Aspergillus nidulans triose phosphate isomerase.

The control sequence may also be a polyadenylation sequence, a sequence which is operably linked to the 3' terminus of the nucleic acid sequence and which, when transcribed, is recognized by the filamentous fungal cell as a signal to add polyadenosine residues to transcribed mRNA. Any polyadenylation sequence, which is functional in the cell, may be used in the present invention.

Preferred polyadenylation sequences for filamentous fungal cells are obtained from the genes encoding Aspergillus oryzae TAKA amylase, Aspergillus niger glucoamylase, Aspergillus nidulans anthranilate synthase, and Aspergillus niger alphaglucosidase.

The control sequence may also be a signal peptide-coding 20 region, which codes for an amino acid sequence linked to the amino terminus of the polypeptide, which can direct the encoded polypeptide into the cell's secretory pathway. The 5' end of the coding sequence of the nucleic acid sequence may inherently 25 contain a signal peptide-coding region naturally linked in translation reading frame with the segment of the coding region, which encodes the secreted polypeptide. Alternatively, the 5' end of the coding sequence may contain a signal peptidecoding region, which is foreign to the coding sequence. 30 foreign signal peptide-coding region may be required where the coding sequence does not normally contain a signal peptidecoding region. Alternatively, the foreign signal peptidecoding region may simply replace the natural signal peptidecoding region in order to obtain enhanced secretion of the

18

polypeptide. The signal peptide-coding region may be obtained from a glucoamylase or an amylase gene from an Aspergillus species, or a lipase or proteinase gene from a Rhizomucor species. However, any signal peptide-coding region, which directs the expressed polypeptide into the secretory pathway of a filamentous fungal cell, may be used in the present invention.

An effective signal peptide coding region for filamentous fungal cells is the signal peptide coding region obtained from the Aspergillus oryzae TAKA amylase gene, Aspergillus niger neutral amylase gene, Rhizomucor miehei aspartic proteinase gene, or Humicola lanuginosa cellulase gene.

The control sequence may also be a propeptide coding region, which codes for an amino acid sequence positioned at the amino terminus of a polypeptide. The resultant polypeptide is known as a proenzyme or propolypeptide (or a zymogen in some cases). A propolypeptide is generally inactive and can be converted to a mature active polypeptide by catalytic or autocatalytic cleavage of the propeptide from the propolypeptide. The propeptide coding region may be obtained from the Rhizomucor miehei aspartic proteinase gene, or the Myceliophthora thermophila laccase gene (WO 95/33836).

Where both signal peptide and propeptide regions are present at the amino terminus of a polypeptide, the propeptide region 25 is positioned next to the amino terminus of a polypeptide and the signal peptide region is positioned next to the amino terminus of the propeptide region.

Expression Vectors

The present invention also relates to recombinant expression vectors comprising a nucleic acid sequence of the present invention, a promoter, and transcriptional and translational stop signals. The various nucleic acid and control sequences described above may be joined together to produce a recombinant

19

expression vector which may include one or more convenient restriction sites to allow for insertion or substitution of the nucleic acid sequence encoding the polypeptide at such sites. Alternatively, the nucleic acid sequence encoding the polypeptide may be expressed by inserting the sequence or a nucleic acid construct comprising the sequence into an appropriate vector for expression. In creating the expression vector, the coding sequence is located in the vector so that the coding sequence is operably linked with the appropriate control sequences for expression, and possibly secretion.

The recombinant expression vector may be any vector (e.g., a plasmid or virus), which can be conveniently subjected to recombinant DNA procedures and can bring about the expression of the nucleic acid sequence encoding the polypeptide. 15 choice of the vector will typically depend on the compatibility of the vector with the filamentous fungal cell into which the vector is to be introduced. The vectors may be linear or closed circular plasmids. The vector may be an autonomously replicating vector, i.e., a vector, which exists as 20 extrachromosomal entity, the replication of which is independent of chromosomal replication, e.g., a plasmid, an extrachromosomal element, a minichromosome, or an artificial chromosome. The vector may contain any means for assuring self-replication. Alternatively, the vector may be one which, 25 when introduced into the filamentous fungal cell, is integrated into the genome and replicated together with the chromosome(s) into which it has been integrated. The vector system may be a single vector or plasmid or two or more vectors or plasmids, which together contain the total DNA to be introduced into the 30 genome of the filamentous fungal cell, or a transposon.

The vectors preferably contain one or more selectable markers, which permit easy selection of transformed cells. A selectable marker is a gene the product of which provides for biocide or viral resistance, resistance to heavy metals,

20

prototrophy to auxotrophs, and the like. A selectable marker for use in a filamentous fungal cell may be selected from the group including, but not limited to, amdS (acetamidase), argB carbamoyltransferase), bar (ornithine (phosphinothricin 5 acetyltransferase), hygB (hygromycin phosphotransferase), niaD (nitrate reductase), pyrG (orotidine-5'-phosphate decarboxylase), sC (sulfate adenyltransferase), (anthranilate synthase), as well as equivalents from other species. Preferred for use in an Aspergillus cell are the amdS 10 and pyrG genes of Aspergillus nidulans or Aspergillus oryzae and the bar gene of Streptomyces hygroscopicus.

The vectors preferably contain an element(s) that permits stable integration of the vector into the host cell genome or autonomous replication of the vector in the cell independent of the genome of the cell.

Host Cells

Another aspect of the present invention relates to host cells comprising a nucleic acid construct or an expression vector of the present invention.

The choice of a host cell in the methods of the present invention will to a large extent depend upon the source of the nucleic acid sequence encoding the polypeptide of interest.

The introduction of an expression vector or a nucleic acid construct into a filamentous fungal cell may involve a process consisting of protoplast formation, transformation of the protoplasts, and regeneration of the cell wall in a manner known per se. Suitable procedures for transformation of Aspergillus cells are described in EP 238 023 and Yelton et al., 1984, Proceedings of the National Academy of Sciences USA 81: 1470-1474. A suitable method of transforming Fusarium species is described by Malardier et al., 1989, Gene 78: 147-156 or in WO 96/00787.

21

"Introduction" means introducing a vector comprising the nucleic acid sequence encoding the polypeptide into a filamentous fungal cell so that the vector is maintained as a chromosomal integrant or as a self-replicating extra5 chromosomal vector. Integration is generally considered to be an advantage as the nucleic acid sequence is more likely to be stably maintained in the cell. Integration of the vector into the chromosome occurs by homologous recombination, non-homologous recombination, or transposition.

For integration into the host cell genome, the vector may 10 rely on the nucleic acid sequence encoding the polypeptide or any other element of the vector for stable integration of the vector into the genome by homologous or nonhomologous recombination. Alternatively, the vector may contain 15 additional nucleic acid sequences for directing integration by homologous recombination into the genome of the host cell. The additional nucleic acid sequences enable the vector to be integrated into the host cell genome at a precise location(s) in the chromosome(s). To increase the likelihood of 20 integration at a precise location, the integrational elements should preferably contain a sufficient number of nucleic acids, such as 100 to 1,500 base pairs, preferably 400 to 1,500 base pairs, and most preferably 800 to 1,500 base pairs, which are highly homologous with the corresponding target sequence to 25 enhance the probability of homologous recombination. integrational elements may be any sequence that is homologous with the target sequence in the genome of the host cell. nucleic acid sequences may be any sequence that is homologous with a target sequence in the genome of the host cell, and, 30 furthermore, may be non-encoding or encoding sequences.

For autonomous replication, the vector may further comprise an origin of replication enabling the vector to replicate autonomously in the host cell.

22

The procedures used to ligate the elements described above to construct the recombinant expression vectors are well known to one skilled in the art (see, e.g., Sambrook, et al., supra).

In a preferred embodiment, the filamentous fungal host cell is a cell of a species of, but not limited to, Acremonium, Aspergillus, Fusarium, Humicola, Mucor, Myceliophthora, Neurospora, Penicillium, Thielavia, Tolypocladium, or Trichoderma.

In a more preferred embodiment, the filamentous fungal cell 10 is an Aspergillus cell. In another more preferred embodiment, the filamentous fungal cell is an Acremonium cell. more preferred embodiment, the filamentous fungal cell is a Fusarium cell. In another more preferred embodiment, the filamentous fungal cell is a Humicola cell. In another more 15 preferred embodiment, the filamentous fungal cell is a Mucor In another more preferred embodiment, the filamentous fungal cell is a Myceliophthora cell. In another more preferred embodiment, the filamentous fungal cell is a Neurospora cell. In another more preferred embodiment, the 20 filamentous fungal cell is a Penicillium cell. In another more preferred embodiment, the filamentous fungal cell is a Thielavia cell. In another more preferred embodiment, the filamentous fungal cell is a Tolypocladium cell. In another more preferred embodiment, the filamentous fungal cell is a 25 Trichoderma cell.

In a most preferred embodiment, the filamentous fungal cell is an Aspergillus awamori, Aspergillus foetidus, Aspergillus japonicus, Aspergillus nidulans, Aspergillus niger or Aspergillus oryzae cell. In another most preferred embodiment, the filamentous fungal cell is a Fusarium bactridioides, Fusarium cerealis, Fusarium crookwellense, Fusarium culmorum, Fusarium graminearum, Fusarium graminum, Fusarium heterosporum, Fusarium negundi, Fusarium oxysporum, Fusarium reticulatum, Fusarium roseum, Fusarium sambucinum, Fusarium sarcochroum,

23

Fusarium sulphureum, Fusarium toruloseum, Fusarium trichothecioides, or Fusarium venenatum cell. In an even most preferred embodiment, the filamentous fungal cell is a Fusarium venenatum (Nirenberg sp. nov.). In another most preferred 5 embodiment, the filamentous fungal cell is a Humicola insolens or Humicola lanuginosa cell. In another most preferred embodiment, the filamentous fungal cell is a Mucor miehei cell. In another most preferred embodiment, the filamentous fungal cell is a Myceliophthora thermophilum cell. In another most 10 preferred embodiment, the filamentous fungal cell Neurospora crassa cell. In another most preferred embodiment, the filamentous fungal cell is a Penicillium purpurogenum cell. In another most preferred embodiment, the filamentous fungal cell is a Thielavia terrestris cell. In another most preferred 15 embodiment, the Trichoderma cell is a Trichoderma harzianum, Trichoderma koningii, Trichoderma longibrachiatum, Trichoderma reesei or Trichoderma viride cell.

Polypeptides having Transcriptional Activation Activity

- Another aspect of the present invention relates to an isolated polypeptide selected from the group consisting of:
- (a) a polypeptide which is encoded in a nucleic acid sequence which hybridizes under low stringency conditions with (i) the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48; (ii) its complementary strand, or (iii) a 25 subsequence of SEQ ID NO:1 or SEQ ID NO: 48 which encodes a polypeptide fragment which has transcriptional activation activity, wherein the low stringency conditions defined are by prehybridization 30 hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micro g/ml sheared and denatured salmon sperm DNA, and 25% formamide, and wash conditions are defined at 50°C for 30 minutes in 2X SSC, 0.2% SDS;

24

- (b) a polypeptide having an amino acid sequence which has at least 50% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49;
- (c) an allelic variant of (a) or (b);
- 5 (d) a fragment of (a), (b), or (c), wherein the fragment has transcriptional activation activity; and
 - (e) a polypeptide comprising the amino acid sequence of SEQ ID NO:3, or an allelic variant thereof.

The transcriptional activator may be isolated using techniques as described herein. As defined herein, an "isolated" polypeptide is a polypeptide which is essentially free of other polypeptides, e.g., at least about 20% pure, preferably at least about 40% pure, more preferably about 60% pure, even more preferably about 80% pure, most preferably about 90% pure, and even most preferably about 95% pure, as determined by SDS-PAGE.

The present invention also relates to isolated polypeptides having an amino acid sequence which has a degree of identity to the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49 of at least about 50%, preferably at least about 55%, preferably at least about 60%, preferably at least about 65%, preferably at least about 70%, preferably at least about 75%, preferably at least about 80%, more preferably at least about 85%, even more preferably at least about 90%, most preferably at least about 95%, and even most preferably at least about 97%, even more preferred at least 99%, which have transcriptional activation activity.

In more preferred embodiment, the transcriptional activator of the present invention comprises the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49 or a fragment thereof, wherein the fragment retains transcriptional activation activity. In a most preferred embodiment, the polypeptide has the amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 49. A fragment of SEQ ID NO: 2 or SEQID NO: 49 is a polypeptide having one or more

WO 01/68864

amino acids deleted from the amino and/or carboxy terminus of this amino acid sequence. Preferably, a fragment of SEQ ID NO:2 or SEQ ID NO: 49 contains at least the polypeptide sequence shown in SEQ ID NO:3.

- The amino acid sequences of the homologous polypeptides may differ from the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49 or SEQ ID NO:3 by an insertion or deletion of one or more amino acid residues and/or the substitution of one or more amino acid residues by different amino acid residues.
- 10 Preferably, amino acid changes are of a minor nature, that is conservative amino acid substitutions that do not significantly affect the folding and/or activity of the protein; small deletions, typically of one to about 30 amino acids; small amino- or carboxyl-terminal extensions, such as an amino-
- 15 terminal methionine residue; a small linker peptide of up to about 20-25 residues; or a small extension that facilitates purification by changing net charge or another function, such as a poly-histidine tract, an antigenic epitope or a binding domain.
- Examples of conservative substitutions are within the group of basic amino acids (such as arginine, lysine and histidine), acidic amino acids (such as glutamic acid and aspartic acid), polar amino acids (such as glutamine and asparagine), hydrophobic amino acids (such as leucine, isoleucine and
- valine), aromatic amino acids (such as phenylalanine, tryptophan and tyrosine), and small amino acids (such as glycine, alanine, serine, threonine and methionine). Amino acid substitutions, which do not generally alter the specific activity are known in the art and are described, for example,
- by H. Neurath and R.L. Hill (1979. The Proteins, Academic Press, New York). The most commonly occurring exchanges are Ala/Ser, Val/Ile, Asp/Glu, Thr/Ser, Ala/Gly, Ala/Thr, Ser/Asn, Ala/Val, Ser/Gly, Tyr/Phe, Ala/Pro, Lys/Arg, Asp/Asn, Leu/Ile, Leu/Val, Ala/Glu, and Asp/Gly as well as these in reverse.

26

In a more preferred embodiment, a transcriptional activator of the present invention is obtained from an Aspergillus niger strain, more preferably from Aspergillus niger AB4.1 (van Haringsveldt, W., et al., 1987. Mol. Gen. Genet. 206:71-75), and most preferably from Aspergillus niger 13PAP2, which has been deposited at DSM as DSM 12298, or a mutant strain thereof, harbouring, e.g., the polypeptide with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49 or SEQ ID NO:3.

In another preferred embodiment, the transcriptional activator of the present invention is the polypeptide encoded in the nucleic acid sequence contained in plasmid pEES, which is contained in *Escherichia coli* DSM 12294 or the nucleic acid sequence shown in SEQ ID NO: 48 encoding the polypeptide shown in SEQ ID NO: 49.

The present invention further relates to methods for producing the transcriptional activator of the present invention comprising (a) cultivating a host cell harbouring a nucleic acid construct or an expression vector comprising a nucleic acid sequence encoding the transcriptional activator of the invention under conditions conducive for production of the polypeptide; and (b) recovering the polypeptide.

Host Cells having Altered Transcriptional Activation Activity

Another aspect of the present invention relates to a host cell which is a mutant of a parent fungal cell useful for the production of a polypeptide in which the parent cell comprises one or more nucleic acid sequences encoding a protease, the transcription of which is activated by a transcriptional activator of the present invention, and the mutant cell produces less of the transcriptional activator and the protease(s) than the parent cell when cultured under the same conditions.

27

The mutant cell may be constructed using methods well known in the art; for example, by one or more nucleotide insertions or deletions of the gene encoding the transcriptional activator.

In a preferred embodiment the mutant cell is obtained by 5 modification or inactivation of a nucleic acid sequence present in the cell and necessary for expression of the transcriptional activator.

In a more preferred embodiment, the nucleic acid sequence is selected from the group consisting of: (a) a nucleic acid sequence having at least 70% identity with the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48; (b) a nucleic acid sequence encoding a polypeptide having an amino acid sequence which has at least 50% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49; (c) a nucleic acid sequence which hybridizes under low stringency conditions with (i) the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48, or (ii) its complementary strand, (d) an allelic variant of (a), (b), or (c); (e) a subsequence of (a), (b), (c), or (d), wherein the subsequence encodes a polypeptide fragment which has transcriptional activation activity; and (f) a subsequence of (a), (b) (c), or (d), wherein the subsequence encodes a polypeptide with the amino acid sequence of SEQ ID NO:3.

In another preferred embodiment the reduced expression of the transcriptional activator in the mutant cell is obtained by modification or inactivation of a control sequence required for the expression of the transcriptional activator. The term "control sequence" is defined, supra, in the section entitled "Nucleic Acid Constructs." In a more preferred embodiment the control sequence in the mutant cell is a promoter sequence or a functional part thereof, i.e., a part, which is sufficient for affecting expression of the nucleic acid sequence. Other control sequences for possible modification include, but are not limited to, a leader, a polyadenylation sequence, a

28

propeptide sequence, a signal sequence, and a transcription terminator.

Modification or inactivation of the gene may be performed by subjecting the parent cell to mutagenesis and selecting for 5 mutant cells in which the capability to produce transcriptional activator has been reduced. The mutagenesis, which may be specific or random, may be performed, for example, by use of a suitable physical or chemical mutagenizing agent, by use of a suitable oligonucleotide, or by subjecting the DNA 10 sequence to PCR generated mutagenesis. Furthermore, the mutagenesis may be performed by use of any combination of these mutagenizing agents.

Examples of a physical or chemical mutagenizing agent suitable for the present purpose include ultraviolet (UV) irradiation, hydroxylamine, N-methyl-N'-nitro-N-nitrosoguanidine (MNNG), O-methyl hydroxylamine, nitrous acid, ethyl methane sulphonate (EMS), sodium bisulphite, formic acid, and nucleotide analogues.

When such agents are used, the mutagenesis is typically performed by incubating the parent cell to be mutagenized in the presence of the mutagenizing agent of choice under suitable conditions, and selecting for mutant cells exhibiting reduced expression of the gene.

Modification or inactivation of the gene may be accomplished by introduction, substitution, or removal of one or more nucleotides in the gene's nucleic acid sequence or a regulatory element required for the transcription or translation thereof. For example, nucleotides may be inserted or removed so as to result in the introduction of a stop codon, the removal of the start codon, or a change of the open reading frame. Such modification or inactivation may be accomplished by site-directed mutagenesis or PCR generated mutagenesis in accordance with methods known in the art. Although, in principle, the modification may be performed in vivo, i.e., directly on the

29

fungal cell expressing the gene to be modified, it is preferred that the modification be performed *in vitro* as exemplified below.

An example of a convenient way to inactivate or reduce sexpression of the gene by a fungal cell of choice is based on techniques of gene replacement or gene interruption. For example, in the gene interruption method, a nucleic acid sequence corresponding to the endogenous gene or gene fragment of interest is mutagenized in vitro to produce a defective nucleic acid sequence which is then transformed into the parent cell to produce a defective gene. By homologous recombination, the defective nucleic acid sequence replaces the endogenous gene or gene fragment. It may be desirable that the defective gene or gene fragment also encodes a marker, which may be used for selection of transformants in which the nucleic acid sequence has been modified or destroyed.

Alternatively, modification or inactivation of the gene may be performed by established anti-sense techniques using a nucleotide sequence complementary to the nucleic acid sequence of the gene. More specifically, expression of the gene by a filamentous fungal cell may be reduced or eliminated by introducing a nucleotide sequence complementary to the nucleic acid sequence, which may be transcribed in the cell and is capable of hybridizing to the mRNA produced in the cell. Under conditions allowing the complementary anti-sense nucleotide sequence to hybridize to the mRNA, the amount of protein translated is thus reduced or eliminated.

A nucleic acid sequence complementary to the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48 may be obtained from any microbial source. The preferred sources are fungal sources, e.g., yeast and filamentous fungi as described supra. Preferred filamentous fungal sources include, but are not limited to, species of Acremonium, Aspergillus, Fusarium, Humicola, Myceliophthora, Mucor, Neurospora, Penicillium,

30

Phanerochaete, Thielavia, Tolypocladium, and Trichoderma.

Preferred yeast sources include, but are not limited to, species of Candida, Hansenula, Kluyveromyces, Pichia, Saccharomyces, Schizosaccharomyces, and Yarrowia. Furthermore, the nucleic acid sequence may be native to the filamentous fungal cell.

In another preferred embodiment, the parent cell harbours a gene having a nucleic acid sequence encoding a polypeptide with an amino acid sequence which has at least 50% identity with the 10 amino acid sequence of SEQ ID NO:2 or SEQID NO: 49.

In another preferred embodiment, the parent cell harbours a gene having a nucleic acid sequence with at least 70% identity with the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48.

In another preferred embodiment, the mutant cell harbours a nucleic acid sequence, which has been modified or inactivated by any of the methods described above and produces less of a protease or a combination of proteases than the parent cell when cultured under identical conditions. The mutant cell produces preferably at least about 25% less, more preferably at least 20 about 50% less, even more preferably at least about 75% less, and even more preferably at least about 95% less of a protease or a combination of proteases than the parent cell when cultured under identical conditions.

In an even more preferred embodiment, the mutant cell produces essentially undetectable amounts of a protease or combination of proteases than the parent cell when cultured under identical conditions.

The protease(s) may be assayed using known methods. In one such method, an aliquot of a 48 hour culture media is incubated 30 with ³H-labelled sperm whale myoglobin at pH 4.0 and the radioactivity in the TCA-soluble fraction is measured (van Noort, J.M., et al., 1991. Anal. Biochem 198:385-390). Other methods have been described for identifying, e.g., aspartic proteinase A. of A. niger (Takahashi, K., 1991. Meth. in

31

Enzymol. 248:146-155), endopeptidases (Morihara, K., 1995. Meth. in Enzymol. 248:242-253), carboxypeptidases (Reminton, J., and Breddam, K., 1994. Meth. in Enzymol. 244:231-248), dipeptidyl peptidase (Ikehara, Y., et al., 244:215-227), and saminopeptidases (Little, G., et al., 1976. Meth. in Enzymol. 45:495-503).

In another preferred embodiment, the mutant cell harbours at least one copy of a nucleic acid sequence encoding a polypeptide of interest.

Another aspect of the present invention relates to a host cell useful for the production of a polypeptide wherein the host cell is a mutant of a parent fungal cell in which the mutant (a) produces more of the transcriptional activator of the present invention as compared to the parent cell when cultured under the same conditions; and (b) comprises a DNA sequence encoding the polypeptide, the transcription of which is activated by the transcriptional activator.

In a preferred embodiment, the host cell produces more of the transcriptional activator than the parent cell when cultured under the same conditions by introducing into the parent cell one or more copies of (i) a nucleic acid sequence encoding a transcriptional activator, (ii) a nucleic acid construct comprising a nucleic acid sequence encoding a transcriptional activator, or (iii) an expression vector as defined above in the section "Expression Vectors".

The nucleic acid construct comprising a nucleic acid sequence encoding a transcriptional activator of the present invention may also comprise one or more nucleic acid sequences which encode one or more factors that are advantageous for directing the expression of the polypeptide, e.g., a transcriptional activator (e.g., a trans-acting factor), a chaperone, and a processing protease. Any factor that is functional in the filamentous fungal cell of choice may be used in the present invention. The nucleic acids encoding one or

more of these factors are not necessarily in tandem with the nucleic acid sequence encoding the polypeptide.

An activator is a protein, which activates transcription of a nucleic acid sequence encoding a polypeptide (Kudla et al., 1990, EMBO Journal 9: 1355-1364; Jarai and Buxton, 1994, Current Genetics 26: 2238-244; Verdier, 1990, Yeast 6: 271-297). The nucleic acid sequence encoding an activator may be obtained from the genes encoding Saccharomyces cerevisiae heme activator protein 1 (hap1), Saccharomyces cerevisiae galactose metabolizing protein 4 (gal4), Aspergillus nidulans ammonia regulation protein (areA), and Aspergillus oryzae alpha-amylase activator (amyR). For further examples, see Verdier, 1990, supra and MacKenzie et al., 1993, Journal of General Microbiology 139: 2295-2307.

- A chaperone is a protein which assists another polypeptide in folding properly (Hartl et al., 1994, TIBS 19: 20-25; Bergeron et al., 1994, TIBS 19: 124-128; Demolder et al., 1994, Journal of Biotechnology 32: 179-189; Craig, 1993, Science 260: 1902-1903; Gething and Sambrook, 1992, Nature 355: 33-45; Puig and Gilbert, 1994, Journal of Biological Chemistry 269: 7764-7771; Wang and Tsou, 1993, The FASEB Journal 7: 1515-11157; Robinson et al., 1994, Bio/Technology 1: 381-384; Jacobs et al., 1993, Molecular Microbiology 8: 957-966). The nucleic acid sequence encoding a chaperone may be obtained from the genes encoding Aspergillus oryzae protein disulphide isomerase or Saccharomyces cerevisiae calnexin, Saccharomyces cerevisiae BiP/GRP78, and Saccharomyces cerevisiae Hsp70. For further examples, see Gething and Sambrook, 1992, supra, and Hartl et al., 1994, supra.
- A processing protease is a protease that cleaves a propeptide to generate a mature biochemically active polypeptide (Enderlin and Ogrydziak, 1994, Yeast 10: 67-79; Fuller et al., 1989, Proceedings of the National Academy of Sciences USA 86: 1434-1438; Julius et al., 1984, Cell 37: 1075-

1089; Julius et al., 1983, Cell 32: 839-852; U.S. Patent No. 5,702,934). The nucleic acid sequence encoding a processing protease may be obtained from the genes encoding Saccharomyces cerevisiae dipeptidylaminopeptidase, Saccharomyces cerevisiae Kex2, Yarrowia lipolytica dibasic processing endoprotease (xpr6), and Fusarium oxysporum metalloprotease (p45 gene).

In a more preferred embodiment, the nucleic acid sequence encoding the transcriptional activator is operably linked to a promoter, or a functional part thereof, which is stronger than the corresponding promoter of the parent cell. In an even more preferred embodiment, the promoter, or a functional part thereof, mediates the expression of a gene encoding an extracellular protease, such as the Aspergillus oryzae alkaline protease, A. oryzae neutral metalloprotease, A. niger aspergillopepsin protease, Fusarium oxysporum trypsin-like protease or F. venenatum trypsin.

The present invention also relates to a host cell useful for the production of a polypeptide wherein the host cell is a mutant of a parent fungal cell in which the mutant comprises

- a) a modification or inactivation of a transcriptional activator of the present invention, or a regulatory sequence thereof, and
 - b) (i) an inducible promoter operably linked to a nucleic acid sequence encoding a transcriptional activator of the present invention, and (ii) a promoter sequence to which the transcriptional activator can bind, operably linked to a nucleic acid sequence encoding the polypeptide, wherein (i) and (ii) can be introduced simultaneously or sequentially.

30

25

The inactive form of the transcriptional activator in (a) above is obtained by inactivation or modification of a nucleic acid sequence present in the cell and necessary for the expression of the native transcriptional activator according to

34

any of the methods as disclosed supra. In a preferred embodiment the inactivation or modification is obtained by methods, which include, but are not limited to, one or more nucleotide insertions, deletions or substitutions, specific or random mutagenesis, gene replacement or gene interruption, and anti-sense techniques using a nucleotide sequence complementary to the nucleic acid sequence of the transcriptional activator. In another preferred embodiment, the inactive form of the native transcriptional activator is obtained by inactivation or modification of a control sequence required for the expression of the transcriptional activator.

In another preferred embodiment, the nucleic acid sequence encoding the native transcriptional activator has the sequence set forth in SEQ ID NO:1 or SEQ ID NO: 48. In another preferred embodiment, the transcriptional activator comprises the polypeptide having the amino acid sequence in SEQ ID NO:3.

The inducible promoter sequence in (b) above may be any promoter sequence, or a functional part thereof, wherein the transcription initiation activity of the promoter can be induced according to the fermentation conditions. Preferably, the induction is mediated by a carbon or nitrogen catabolite. In a preferred embodiment, the promoter is the amdS promoter of Aspergillus nidulans or A. oryzae, the niaD promoter of A. nidulans, A. oryzae or A. niger, the niiA promoter of Aspergillus species, the alkaline phosphatase promoter of Aspergillus sp., the acid phosphatase promoter of Aspergillus sp., or the alcA promoter of A. niger.

In another preferred embodiment, the host cell further comprises a promoter sequence, wherein the promoter sequence can be activated by the transcriptional activator and is operably linked to the nucleic acid sequence encoding the polypeptide.

The promoter sequence activated by the transcriptional activator of the present invention may be any promoter

sequence, or a functional part thereof, selected from the group which includes but is not limited to promoters obtained from the genes encoding Aspergillus oryzae TAKA amylase, Rhizomucor aspartic proteinase, Aspergillus niger 5 alpha-amylase, Aspergillus niger acid stable alpha-amylase, Aspergillus niger or Aspergillus awamori glucoamylase (glaA), Rhizomucor miehei lipase, Aspergillus oryzae alkaline protease, Aspergillus oryzae triose phosphate isomerase, Aspergillus nidulans acetamidase, the NA2-tpi promoter (a hybrid of the 10 promoters from the genes encoding Aspergillus niger neutral alpha-amylase and Aspergillus oryzae triose phosphate isomerase), and mutant, truncated, and hybrid promoters thereof. · Particularly preferred promoters for use filamentous fungal cells are a promoter, or a functional part 15 thereof, from a protease gene; e.g., from the Fusarium oxysporum trypsin-like protease gene (U.S. Patent 4,288,627), Aspegillus oryzae alkaline protease gene (alp), Aspergillus niger pacA gene, Aspergillus oryzae alkaline protease gene, A. oryzae neutral metalloprotease gene, A. niger 20 aspergillopepsin protease gene, or F. venenatum trypsin gene.

In another preferred embodiment, the host cell harbours at least one copy of a nucleic acid sequence encoding a polypeptide.

In another preferred embodiment, the host cell, which
25 expresses the transcriptional activator of the present
invention produces less of one or more native proteases than
the parent cell when cultured under identical conditions. The
protease(s) may be assayed using any of the methods described
above. In a more preferred embodiment, an aliquot from a 4830 hour culture media is incubated with ³H-labelled sperm whale
myoglobin at pH 4.0 and the radioactivity in the TCA-soluble
fraction is measured (van Noort, J.M., et al., supra).

The nucleic acid constructs described herein may be introduced into a parent fungal cell according to any of the

36

methods as described supra in the section, "Host Cells" to obtain a host cell useful for the production of a polypeptide. In a preferred embodiment the nucleic acid construct is integrated into the chromosome of the cell. In another preferred embodiment the nucleic acid construct is maintained as a self-replicating extra-chromosomal vector.

It will be understood that the methods of the present invention are not limited to a particular order for obtaining the mutant fungal cell. The modification of the second nucleic acid sequence may be introduced into the parent cell at any step in the construction of the cell for the production of a polypeptide.

Producing a Polypeptide

Another aspect of the present invention relates to methods of producing a polypeptide in a host cell of the present invention, comprising: (a) cultivating the host cell which harbours a gene encoding the polypeptide in a nutrient medium suitable for production of the polypeptide; and (b) recovering the polypeptide from the nutrient medium of the host cell.

In one embodiment, the host cell which is a mutant of a parent fungal cell in which the parent cell comprises one or more nucleic acid sequences encoding a protease, the transcription of which is activated by a transcriptional 25 activator of the present invention, and the mutant cell produces less of the transcriptional activator and the protease(s) than the parent cell when cultured under the same conditions.

In another embodiment, the host cell is a mutant of a parent fungal cell in which the mutant (a) produces more of the transcriptional activator of the present invention as compared to the parent cell when cultured under the same conditions; and (b) comprises a DNA sequence encoding the polypeptide, the

37

transcription of which is activated by the transcriptional activator.

In another embodiment, the host cell is a mutant of a parent fungal cell in which the mutant comprises (a) a modification or inactivation of a transcriptional activator of the present invention or a regulatory sequence thereof, and (b) an inducible promoter operably linked to a nucleic acid sequence encoding a transcriptional activator of the present invention and a promoter sequence to which the transcriptional activator can bind, operably linked to a nucleic acid sequence encoding the polypeptide, wherein (i) and (ii) can be introduced simultaneously or sequentially.

The host cells of the present invention are cultivated in a nutrient medium suitable for production of the polypeptide of 15 interest using methods known in the art. For example, the cells may be cultivated by shake flask cultivation, small-scale or large-scale fermentation (including continuous, batch, fedbatch, or solid state fermentations) in laboratory or industrial fermentors performed in a suitable medium and under 20 conditions allowing the polypeptide to be expressed and/or isolated. The cultivation takes place in a suitable nutrient medium comprising carbon and nitrogen sources and inorganic salts, using procedures known in the art (see, e.g., Bennett, J.W. and LaSure, L., eds., More Gene Manipulations in Fungi, 25 Academic Press, CA, 1991). Suitable media are available from commercial suppliers or may be prepared using published compositions (e.g., in catalogues of the American Type Culture Collection). If the polypeptide is secreted into the nutrient medium, the polypeptide can be recovered directly from the 30 medium. If the polypeptide is not secreted, it is recovered from cell lysates.

The resulting polypeptide may be isolated by methods known in the art. For example, the polypeptide may be isolated from the nutrient medium by conventional procedures including, but

38

not limited to, centrifugation, filtration, extraction, spray evaporation, or precipitation. The isolated polypeptide may then be further purified by a variety of procedures known in the art including, but not limited to, 5 chromatography (e.g., ion exchange, affinity, hydrophobic, chromatofocusing, and size exclusion), electrophoretic (e.g., procedures preparative isoelectric differential solubility (e.g., ammonium sulfate precipitation), or extraction (see, e.g., Protein Purification, J.-C. Janson 10 and Lars Ryden, editors, VCH Publishers, New York, 1989).

The polypeptide may be detected using methods known in the art that are specific for the polypeptide. These detection methods may include use of specific antibodies, formation of an enzyme product, disappearance of an enzyme substrate, or SDS-PAGE. For example, an enzyme assay may be used to determine the activity of the polypeptide. Procedures for determining enzyme activity are known in the art for many enzymes.

In the methods of the present invention, the host cell produces at least about 20% more, preferably at least about 50% 20 more, more preferably at least about 100% more, even more preferably at least about 200% more, and most preferably at least about 300% more of the polypeptide than a corresponding parent cell when cultivated under the same conditions.

The polypeptide may be any polypeptide whether native or heterologous to the mutant filamentous fungal cell. The term "heterologous polypeptide" is defined herein as a polypeptide, which is not produced by a cell. The term "polypeptide" is not meant herein to refer to a specific length of the encoded produce and therefore encompasses peptides, oligopeptides and proteins. The polypeptide may also be a recombinant polypeptide, which is a polypeptide native to a cell, which is encoded by a nucleic acid sequence, which comprises one or more control sequences, foreign to the nucleic acid sequence, which are involved in the production of the polypeptide. The

39

polypeptide may be a wild-type polypeptide or a variant thereof. The polypeptide may also be a hybrid polypeptide, which contains a combination of partial or complete polypeptide sequences obtained from at least two different polypeptides where one or more of the polypeptides may be heterologous to the cell. Polypeptides further include naturally occurring allelic and engineered variations of the above-mentioned polypeptides.

In a preferred embodiment, the polypeptide is an antibody or portions thereof, an antigen, a clotting factor, an enzyme, a hormone or a hormone variant, a receptor or portions thereof, a regulatory protein, a structural protein, a reporter, or a transport protein.

In a more preferred embodiment, the enzyme is an 15 oxidoreductase, transferase, hydrolase, lyase, isomerase, or ligase.

In an even more preferred embodiment, the enzyme is an aminopeptidase, amylase, carbohydrase, carboxypeptidase, catalase, cellulase, chitinase, cutinase, deoxyribonuclease, 20 dextranase, esterase, alpha-galactosidase, beta-galactosidase, glucoamylase, alpha-glucosidase, beta-glucosidase, invertase, laccase, haloperoxidase, lipase, mannosidase, mutanase, oxidase, pectinolytic enzyme, peroxidase, phytase, polyphenoloxidase, proteolytic enzyme, ribonuclease, 25 transglutaminase, or xylanase.

In another even more preferred embodiment, the polypeptide is human insulin or an analog thereof, human growth hormone, erythropoietin, or insulinotropin.

The nucleic acid sequence encoding a heterologous 30 polypeptide may be obtained from any prokaryotic, eukaryotic, or other source. For purposes of the present invention, the term "obtained from" as used herein in connection with a given source shall mean that the polypeptide is produced by the

40

source or by a cell in which a gene from the source has been inserted.

In the methods of the present invention, the mutant filamentous fungal cells may also be used for the recombinant 5 production of polypeptides, which are native to the cell. native polypeptides may be recombinantly produced by, e.g., placing a gene encoding the polypeptide under the control of a different promoter to enhance expression of the polypeptide, to expedite export of a native polypeptide of interest outside the 10 cell by use of a signal sequence, and to increase the copy number of a gene encoding the polypeptide normally produced by the cell. The present invention also encompasses, within the scope of the term "heterologous polypeptide", such recombinant production of polypeptides native to the cell, to the extent 15 that such expression involves the use of genetic elements not native to the cell, or use of native elements which have been manipulated to function in a manner that do not normally occur in the filamentous fungal cell. The techniques used to isolate or clone a nucleic acid sequence encoding a heterologous 20 polypeptide are known in the art and include isolation from genomic DNA, preparation from cDNA, or a combination thereof. The cloning of the nucleic acid sequences from such genomic DNA can be effected, e.g., by using the well known polymerase chain reaction (PCR), see, for example, Innis et al., 1990, PCR 25 Protocols: A Guide to Methods and Application, Academic Press, New York. The cloning procedures may involve excision and isolation of a desired nucleic acid fragment comprising the nucleic acid sequence encoding the polypeptide, insertion of the fragment into a vector molecule, and incorporation of the 30 recombinant vector into the mutant fungal cell where multiple copies or clones of the nucleic acid sequence will be replicated. The nucleic acid sequence may be of genomic, cDNA, RNA, semisynthetic, synthetic origin, or any combinations thereof.

41

In the methods of the present invention, heterologous polypeptides may also include fused or hybrid polypeptides in which another polypeptide is fused at the N-terminus or the Cterminus of the polypeptide or fragment thereof. A fused 5 polypeptide is produced by fusing a nucleic acid sequence (or a portion thereof) encoding one polypeptide to a nucleic acid sequence (or a portion thereof) encoding another polypeptide. Techniques for producing fusion polypeptides are known in the art, and include, ligating the coding sequences encoding the 10 polypeptides so that they are in frame and expression of the fused polypeptide is under control of the same promoter(s) and terminator. The hybrid polypeptides may comprise a combination of partial or complete polypeptide sequences obtained from at least two different polypeptides wherein one or more may be 15 heterologous to the mutant fungal cell. An isolated nucleic acid sequence encoding a heterologous polypeptide of interest may be manipulated in a variety of ways to provide for expression of the polypeptide. Expression will be understood to include any step involved in the production of the 20 polypeptide including, but not limited to, transcription, posttranscriptional modification, translation, post-translational modification, and secretion. Manipulation of the nucleic acid sequence encoding a polypeptide prior to its insertion into a vector may be desirable or necessary depending on the 25 expression vector. The techniques for modifying nucleic acid sequences utilizing cloning methods are well known in the art.

The present invention is further described by the following examples, which should not be construed as limiting the scope of the invention.

42

EXAMPLES

Materials

Chemicals used as buffers and substrates were commercial products of at least reagent grade.

Strains

AB4.1: a strain of Aspergillus niger which is a cspA1 pyrG1 derivative of strain ATTC 9029 (van Hartingsveldt, W., et al., 10 1987. Mol. Gen. Genet. 206:71-75; Bos, C.J., et al., Curr. Genet. 14:437-443)

AB1.13: a protease deficient strain of Aspergillus niger derived from UV mutagenesis of AB4.1 (Mattern, I.E., et al., 1992. Mol. Gen. Genet. 234:332-336)

15 13PAP2: an AB1.13 derivative containing multiple copies of the A. nidulans amdS gene (Corrick, R. A., et al, 1987. Gene 53: 63 - 71) under control of the pepA promoter of A. niger (Jarai G. and Buxton F. 1994. Curr Genet 26:238-244). The strain has a protease deficient phenotype and is unable to grow 20 on medium containing acetamide as the sole nitrogen source. Strain 13PAP2 has been deposited at DSM under the name DSM No. 12298.

4PAP6: an AB4.1 derivative containing multiple copies of the of A. nidulans amdS gene under control of the pepA promoter of 25 A. niger. The strain does not have a protease deficient phenotype and is able to grow on medium containing acetamide as the sole nitrogen source.

N402: a strain of Aspergillus niger, deposited at the ATCC (Manassas VA, USA) as ATCC Number: 64974

MC1046: a strain of *E. coli*, deposited at the ATCC as ATCC Number: 35467

A.oryzae IFO4177: available from Institute for Fermentation, Osaka; 17-25 Juso Hammachi 2-Chome Yodogawa-Ku, Osaka, Japan.

WO 01/68864

43

HowB101: described in WO 97/35956.

Plasmids

pPAP: constructed as described below in Example 1 and 5 shown in Figure 1

pAopyrGcosArp1: constructed as described below in Example 1 and shown in Figure 2

pEES1: constructed as described below in Example 1 and shown in Figure 3

p3SR2: contains the A. nidulans amdS gene as described by C.M. Corrick, A.P. Twomey, and M.J. Hynes (1987. Gene 53: 63-71)

pABPYRG*-Not: contains an inactivated pyrG gene as described by Verdoes, J.C., et al. (1994. Gene 145: 179-187)

pHelpl: contains the *pyrG* gene from A. oryzae as a selective marker and the AMA1 sequences which enable autonomous replication in A. niger, cloned into the E. coli vector pIC20R, as described by Gems, D., et al. (1991. Gene 98: 61-67)

pAnscos1: contains two cos sites as described by 20 Osiewacz, H.D. (1994. Curr. Genet. 26: 87-90)

pAO4-2: contains the A. oryzae pyrG gene as described by De Ruiter-Jacobs, Y.M.J.T., et al. (1989. Curr. Genet. 16: 159-163)

pAO4-13: contains the A. oryzae pyrG gene as described by De 25 Ruiter-Jacobs, Y.M.J.T., et al. (1989. Curr. Genet. 16:159-163)

pUC19: as described by Yanisch-Perron, C., Vieira, J. and Messig, J. (1985, Gene 33:103-119)

pDV8: described in Example 8 and shown in Fig. 7.

pJaL554: described in Example 8 and shown in Fig. 8

Deposit of Biological Materials

The following biological material has been deposited under the terms of the Budapest Treaty with the Deutsche Sammlung von

44

Microorganismen und Zellkulturen GmbH, Mascheroder Weg 1b, D-38124 Braunschweig, Germany and given the following accession number:

Deposit Accession Number Date of Deposit

5 Escherichia coli, pEES DSM 12294 1998-07-14

Aspergillus niger 13PAP2 DSM 12298 1998-07-14

The strain has been deposited under conditions that assure that access to the culture will be available during the pendency of this patent application. The deposit represents a substantially pure culture of the deposited strain. The deposit is available as required by foreign patent laws in countries wherein counterparts of the subject application, or its progeny are filed. However, it should be understood that the availability of a deposit does not constitute a license to practice the subject invention in derogation of patent rights granted by governmental action.

EXAMPLES

20

Example 1

Cloning of the A. niger prtT transcriptional activator

The prtT gene was cloned from 13PAP2, an A. niger mutant strain which is unable to express the amdS gene regulated by the pepA protease gene promoter and has a protease deficient phenotype (prt-).

Construction of the A. niger 13PAP2 reporter strain

The plasmid pPA1 was contructed by ligation of the following 30 three fragments:

- 1) the E. coli vector pBlueScript II SK (Stratagene Cloning Systems, La Jolla CA, USA) digested with EcoRI and KpnI;
- 2) a 1.4 kb EcoRI/BamHI restriction fragment containing the 1.2 kb promoter region of the pepA gene linked to about

45

130 bp of the amdS coding sequence from the start codon to an internal BamHI site, amplified by PCR; and

3) a 2.1 kb BamHI/KpnI fragment from p3SR2 which contains most of the A. nidulans amdS gene

5

Fragment 2 was constructed in two steps. In a first step genomic DNA from A. niger N402 prepared from protoplasts as described below in the section "Construction of the Cosmid Library" was used as the template, and the two oligonucleotides shown below, pepApr and pepA/amdS, were used as primers:

PepApr: CGG AAT TCG CAT GCT GGA GGT GCT TCT AA (SEQ ID NO: 6)

pepA/amdS:TTC CCA GGA TTG AGG CAT TTT GAC CAC GAG AAT (SEQ ID NO:

15

The 1200 bp PCR product obtained from this reaction was then used as a primer in a second PCR reaction together with the oligonucleotide MBL1213 shown below, and plasmid p3SR2 as the template.

20

MBL1213: TAA CTT CCA CCG AGG TC (SEQ ID NO: 8)

The product obtained by ligation of the three fragments described above was subsequently transfected into $E.\ coli\ DH5\alpha.$

In the final construction procedure, pPA1 was digested with NotI and ligated to a 3.8 kb NotI fragment from pABPYRG*-Not, resulting in plasmid pPAP which is shown in Fig. 1. pPAP was transformed into A. niger AB 1.13, and a transformant with pPAP integrated into the pyrG locus in multicopy was isolated. A spontanous 5 flourotic acid (FOA) resistant, uridine-requiring mutant of this transformant that could be complemented with the pyrG gene was named 13PAP2.

46

Construction of pAopyrGcosArp1

The plasmid pAopyrGcosArp1 was constructed by ligation and subsequent transfection into $E.\ coli\ DH5\alpha$ of the following three fragments:

- 1) the E. coli vector pHelp1 cut with Acc65I and BamHI
 - 2) a 3.0 kb BamHI/HindIII fragment from pAnscosl containing two cos sites
 - 3) a 3.2 kb Acc65I/HindIII fragment from pAO4-2 containing the A. oryzae pyrG gene

10

The resulting plasmid, pAopyrGcosArp1, is self-replicating in Aspergilli and can be selected for by growth on medium lacking uridine. pAopyrGcosArp1 is depicted in Fig. 2.

15 Construction of the cosmid library.

A cosmid library of Aspergillus niger was constructed using the "SuperCos1 cosmid vector kit" (Stratagene Cloning Systems, La Jolla CA, USA) according to the supplier's instructions.

Genomic DNA from $A.\ niger$ N402 was prepared from protoplasts 20 made by standard procedures.

After isolation the protoplasts were pelleted by centrifugation at 2000 rpm for 10 minutes in a Beckman GS-6R; the pellet was then suspended in a buffer containing 22.5 mM triisonaphtalene sulphonic acid, 275 mM para-aminosalicylic acid, 25 0.2 M Tris-HCl (pH 8.5), 0.25 M NaCl and 50 mM EDTA immediately followed by addition of 1 volume of phenol/chloroform (1:1). After careful mixing and centrifugation at 3000 rpm for 20 minutes the aqueous phase was decanted and DNA was precipitated using standard procedures.

The size of the genomic DNA was analysed by electrophoresis on a 0.3% agarose gel run for 20 hours at 30 volts at 4°C. The ethidium bromide stained gel showed that the recovered DNA ranged in size from 50 to greater than 100 kb. The DNA was partially digested using MboI. The size of the digested DNA was

47

30 to 50 kb as determined by the same type of gel analysis as above. The pAopyrGcosArpl vector, purified using a kit from QIAGEN (Venlo, The Netherlands) following the manufacturer's instructions, was digested with BamHI, dephosphorylated and gel purified. Ligation and packaging were performed following standard procedures.

After titration of the library, all of the packaging mix from a single ligation and packaging was transfected into the host cell, MC1046, and plated on 50 μ g/ml ampicillin LB plates. 10 Approximately 40,000 colonies were obtained. Cosmid preparations from 10 colonies showed that they all had inserts of the expected size. The 40,000 colonies were then soaked in LB medium and scraped off of the plates, then aliquoted for storage in 15% glycerol at -80°C. This represents an approximate 40-15 fold amplification of the A. niger genome.

Selection of A. niger prtT clones

Cosmid DNA was prepared from the library and introduced into 13PAP2 according to the transformation procedure described by P.J. Punt and C.A.M.J.J. Van Den Hondel (1992. Methods Enzymol 216: 447-457). Repeated efforts to select for the pyrG marker only resulted in a recovery of between 4000 to 30,000 transformants. A double selection for the pyrG marker and growth on medium containing acetamide as the sole nitrogen source resulted in a total of 65 primary transformants from five different experiments.

Each primary transformant was screened for protease activity, growth on medium containing acetamide as the sole nitrogen source and instability of the these two characteristics. An acetamidase phenotype, screened by growth on medium containing acetamide, is an indication of acetamidase activity resulting from activation of the pepA promoter in the reporter cassette in which the pepA promoter is linked to the amdS coding sequence. A protease phenotype was screened using minimal me-

48

dium plates containing dialyzed skim milk as the sole nitrogen source (Mattern, I.E., et al., 1992. Mol Gen Genet 234:332-336). On these plates the wild-type AB4.1 strain makes a clear halo whereas the AB1.13 mutant produces a very small halo. This difference is not due to differences in the activity of pepA since a pepA deleted strain can also produce a large halo on these plates. Therefore, a large halo on milk plates indicates activation of other extracellular proteases.

Instability was tested by growing diluted spore stocks on a medium containing uridine. Single-spore-derived colonies were picked from these plates and tested for protease activity and growth on acetamide. The screening results revealed that in more than 70% of the colonies both characteristics were lost. Therefore, the two phenotypes were either lost or retained together, indicating that activation of the pepA promoter and other protease promoters is coordinately regulated and linked to the presence of the pyrG marker. The gene responsible for this phenotype was named prtT. Twelve acetamidase[†], protease[†] transformants were then isolated.

20

Isolation of the A.niger prtT gene

In order to rescue the prtT gene from the acetamidase*, protease* transformants of 13PAP2, DNA was prepared from mycelium grown in minimal medium as previously described. This DNA was used in an attempt to transform competent E. coli DH5α cells. Several hundreds of ampicillin-resistant colonies were obtained. DNA analysis showed they all contained sequences derived from the pHelp1 plasmid. Cosmid DNA isolated from E. coli colonies was then retransformed into 13PAP2. Two DNA samples gave rise to transformants, which showed both growths on acetamide containing medium and increased protease activity. DNA from one of the cosmids, ACR1, was then digested with several restriction enzymes. The resulting fragments were then co-transformed with pAopyrGcosArp1 into strain 13PAP2. EcoRI,

PstI, BamHI and KpnI digestion of ACR1 gave rise to transformants capable of growth on acetamide and high protease activity, whereas SalI and HindIII digests did not. Because EcoRI digestion gave the simplest pattern, separate EcoRI fragments 5 were gel-isolated and with pAopyrGcosArp1 used to co-transform Only one fragment, a 15 kb EcoRI fragment, gave rise to transformants capable of growth on acetamide-containing me-This fragment was subcloned in pBluescript II SK in order to subclone prtT from the cosmid. Since the insert of this 10 clone was still rather large, separate PstI bands were gel isolated and each was co-transformed with pAopyrGcosArpl into 13PAP2. Only one band, a 2.5 kb PstI fragment, gave rise to transformants that could grow on acetamide-containing medium. This fragment was subcloned in pBlueScript II SK. Four sub-15 clones, ClE 0.7, ClE 1.8, NcE 1.1 and NcE 1.4, were constructed from this plasmid based on the restriction map. In addition, a 6.5 kb SstI/EcoRI fragment encompassing the 2.5 kb PstI fragment was subcloned, resulting in pEES1 (shown in Fig. 3).

Southern blot analysis of genomic DNA from AB4.1 showed the 20 presence of only one copy of prtT.

Example 2

Sequencing of the A. niger prtT gene and analysis of the sequence

All sequence reactions were prepared using dRhodamine Terminator Cycle Sequencing Kits or BigDye™ Terminator Cycle Sequencing Kits from the Perkin-Elmer Corporation (Branchburg NJ, USA). The reactions were run on an ABI PRISM® 377 DNA Sequencer (Perkin-Elmer Corporation) following the manufacturer's instructions.

The prtT gene was sequenced from the genomic clones ClE 0.7, ClE 1.8, NcE 1.1, NcE 1.4 and pEES1. The sequence specific primers used are listed below:

122958: CGA TCG ATG ACT GCC TGT (SEQ ID NO: 9)

50

```
122956:
            AGA GAC ACA TAG TGC CTT (SEQ ID NO: 10)
  122959:
            GCT TAT AGT CGA TAG CGC (SEQ ID NO: 11)
  122960:
            CCT CTC TCC AGC GAT GGT (SEQ ID NO: 12)
            ATG GAA TAC ATA CTG CTT (SEQ ID NO: 13)
  122962:
5 122961:
            ATG AAA CCC ACT GTA GCT (SEQ ID NO: 14)
            TGC TCG ATA AGC GGG TCC (SEQ ID NO: 15)
  122963:
  122964:
            AAT CTT ATG GAC CCG CTT (SEQ ID NO: 16)
  124289:
            CCC CGG GAA ACA AGA ACA GG (SEQ ID NO: 17)
  124290:
            GTT GGC GGA CCT TGA CTA TG (SEQ ID NO: 18)
10 125112:
            ACA GCT ACA GTG GGT TTC ATC T (SEQ ID NO: 19)
  125111:
            AGT CAA CGG GGG AAG TCT C (SEQ ID NO: 20)
  128330:
            CTA GCA GCG TAT CGG TCA GC (SEQ ID NO: 21)
  130887:
            CTT GGA AAA GAA ACG ATA G (SEQ ID NO: 22)
  130888:
            AAC GTA CGC TTT CCT CCT T (SEQ ID NO: 23)
            GGG TCC GTC CAG TCC GTT CTT (SEQ ID NO: 24)
15 134135:
```

-48 reverse: AGC GGA TAA CAA TTT CAC ACA GGA (SEQ ID NO: 25)

-40 universal: GTT TTC CCA GTC ACG AC (SEQ ID NO: 26)

20

A mutant allele of the gene was obtained by PCR amplification of genomic DNA isolated from the mutant strain AB1.13 using the following primers:

25 PstI: TC ATC CCT GGT GTT ACT GC (SEQ ID NO: 27)
PstII: C ATG GAT TGG CTG GCC G (SEQ ID NO: 28)

The complete DNA sequence of the prtT gene is shown in SEQ ID NO:1. The sequence of the PCR fragment of the mutant allele 30 is shown in SEQ ID NO:4.

Analysing the DNA sequence SEQ ID NO:1 using the computer software Netgene 2 (S.M. Hebsgaard, P.G. Korning, N. Tolstrup, J. Engelbrecht, P. Rouze, S. Brunak (1996. Nucleic Acids Re-

51

search 24: 3439-3452) suggested the existence of 5 exons (see annotations to SEQ ID NO 1).

Analysis of the A. niger prtT cDNA

- mRNA was purified from total RNA (isolated according to the DNA isolation method described above in Example 1) using a commercial poly(A)⁺ RNA isolation kit (Pharmacia, Uppsala SE) from a culture of A. niger grown under conditions favourable for protease production (J.P.T.W. Van Den Hombergh, et al., 1997.
- 10 Eur. J. Biochem. 247:605-613). Double stranded cDNA was prepared using standard procedures and used for PCR reactions with the following primers:

oligo-dT primer: T20N

15 Prt270n: TACTCTCCAGATTGCCTG (SEQ ID NO: 29)

Prt1420r: TGAGATACCACTCAGCAG (SEQ ID NO: 30)
prt1350n: TGCACTTCTCTGTCTCTG (SEQ ID NO: 31)
Prt2365r: GACTTCTGGCATCAGTTG (SEQ ID NO: 32)

prt2320n: CTCATGGATGGCATGATC (SEQ ID NO: 33)

20

A PCR reaction with the primers Prt270n and Prt1420r produced a fragment of approximately 1.0 kb. The fragment was cloned into a pGEM-T vector (Promega Corp., Madison WI, USA), and the insert in the resulting plasmid was sequenced using the primers 122958, 122960, -40 universal and -48 reverse. The result confirmed the presence of two introns in this part of the gene.

A second PCR reaction with the primers Prt1350n and Prt2365r produced a fragment of approximately 0.9 kb. This fragment was also cloned in a pGEM-T vector, and the insert in the resulting plasmid was sequenced using the primers 124289, 124290, -40 universal and -48 reverse. The result confirmed the presence of a single intron in this part of the gene.

52

Another PCR reaction with the oligo-dT primer and primer Prt2320n produced a fragment of approximately 350 bp. This fragment was also cloned in a pGEM-T vector. Sequencing of the insert using primers -40 universal and -48 reverse showed that the fragment contained the 3' part of prtT and confirmed the presence of another intron.

The deduced protein sequence of the translated prtT gene is shown in SEQ ID NO:2. The deduced protein sequence of the translated mutant allele prt13 is shown in SEQ ID NO:5. A com10 parison of SEQ ID NO:2 and SEQ ID NO:5 indicates that the only difference between the two is in position 112 where the leucine residue in the translated prtT gene is replaced by proline in the translated prt13 gene.

Analysis of the deduced PrtT protein sequence reveals the

15 presence of a Zinc(II)2Cys6 binuclear cluster DNA binding motif

(SEQ ID NO:2, residues 47-81). This motif defines the GAL4

class of fungal transcriptional activators (Reece, M. J., and

Ptashne, M. 1993. Science 261: 909-911). The presence of the

motif in the prtT gene strongly indicates that prtT is a tran
20 scriptional activator.

EXAMPLE 3:

Disruption of the prtT gene in a wild-type A. niger strain

A plasmid was constructed in which the upstream and downstream sequences of the prtT gene are separated by the A.
oryzae pyrG gene. Plasmid pEES1 was digested with MunI and
NheI, which removed a 2.1 kb fragment containing most of the
coding sequence of prtT. A 2.3 kb EcoRI/NheI fragment from
pAO4-13 containing the A. oryzae pyrG gene was cloned in the
MunI and NheI sites of pEES1. The resulting plasmid, shown in
Fig. 4, was named pDprt. This construct was then used to
transform A. niger strain AB4.1 to uridine prototrophy. About
150 uridine prototrophic transformants were then analyzed for
protease activity on skim milk containing plates. Five of

53

these did not make a halo on these plates indicating that protease activity was very low. Comparison of strains with a disrupted prtT gene and the mutant AB1.13 strain did not show any differences in protease activity or phenotype.

5

Example 4

Overexpression of A. niger PrtT

A plasmid, pGPprt, (Figure 5) containing the coding region and 3' noncoding sequences of prtT fused to the promoter of the A. niger gpd gene was constructed. The gpd gene codes for glyceraldehyde-3-phosphate dehydrogenase and a constitutively expressed enzyme involved in primary metabolism. The promoter used was a fragment upstream of the coding region.

15 The plasmid is transformed into A. niger AB4.1 by cotransformation with the pyrG selection plasmid pAO4-13. Transformants with increased prtT transcription as determined by Southern blot analysis is analysed for increased protease expression.

20 Example 5

Isolation of the Zn^{2+} -finger from the A. oryzae prtT gene

The A. niger prtT gene is shown in SEQ ID NO: 1. The protein sequence deduced from the DNA sequence of prtT (SEQ ID NO: 2) contains a so called Zn²⁺-finger motif expected to be responsible for the DNA binding of the transcriptional activator encoded by prtT. The Zn²⁺-finger motif has the following amino acid sequence: Met Thr Ala Cys His Thr Cys Arg Lys Leu Lys Thr Arg Cys Asp Leu Asp Pro Arg Gly His Ala Cys Arg Arg Cys Leu Ser Leu Arg Ile Asp Cys (SEQ ID NO: 34).

Degenerate primers able to code for amino acid sequences from the motif were designed and synthesized by DNA Technology A/S, Forskerparken, Gustav Wieds vej 10, DK-8000 Aarhus C, Denmark. The primers had the following sequences:

137396: A T G A C C/T G C C/T T G C/T C A C/T A C C/T T G (SEQ ID NO: 35)

137397: A A/G A/G C A A/G/C/T C G A/G/C/T C G A/G C A A/G G C A/G T G (SEQ ID NO: 36)

- The primers were used in a PCR reaction with genomic A.

 oryzae IFO4177 DNA as template. The reaction was performed in a
 total volume of 100 μl containing 154 pmol of primer 137396 and
 10164 pmol of primer 137397. 30 PCR cycles with 56°C as annealing temperature and 30 seconds elongation time were run. An10 other PCR reaction using A. niger genomic DNA and the primers
 137394: ATGACTGCCTGTCACACATG (SEQ ID NO: 37)
 - and 137395: AGACAGCGACGGCACGCATG (SEQ ID NO:38), which are specific for the A. niger prtT gene, was also run. In this reaction 10 pmol of each primers was used in a 100 μ l reaction.
- After electrophoresis three approximately equally intense bands could be seen in the A. oryzae reaction and two bands in the A. niger reaction. One of the bands in the A. niger reaction was more intense than the other and further had the expected size.
- 20 One of the A. oryzae bands had the same size as the most intense A. niger band and was isolated from the gel. The fragment was cloned into the vector pCR2.1 (Invitrogen™). Plasmids from two individual colonies were sequenced. The sequences are shown in figure 1. The two sequences differ at the end reflecting
- 25 their origin in different degenerate primers. They are identical in the middle 40bp, which are amplified from the genomic DNA. These 40 basepairs encode a polypeptide identical to a part of the Zn²⁺-finger of the A. niger prtT gene.

30 Example 6 .

Isolation of the N-terminal of the A. oryzae prtT gene

The inverse PCR method was used to isolate the A. oryzae prtT gene. The primers 144428: CACCGAGTTTTAAGCTTGCGG (SEQ ID NO: 39) and 144429:GCGATCTTGATCCACGAGGG (SEQ ID NO: 40) were

synthezised by DNA Technology A/S (Denmark). Genomic DNA was cut with a number of restriction enzymes and religated. The ligation mixtures were used as templates in PCR reactions with the primers 144428/144429. In a reaction with BamHI restricted 5 and religated DNA as template a fragment of approximately 2.5kb was observed after electrophoresis on an agarose gel. The fragment was labelled with ^{32}P by the random priming method and used as a probe against a filter containing a gridded cosmid library of genomic A. oryzae DNA. The construction of the li-10 brary is described in WO 98/01470. The cosmid 11F8 showed a positive hybridization signal with the probe. A Southern blot containing DNA from 11F8 and genomic DNA restricted with BamHI, EcoRI, PstI or XhoI was probed with the 2.5kb inverse PCR fragment. The size of hybridizing bands from genomic DNA were com-15 pared with those from the cosmid DNA. Apparently some rearrangement of the cosmid had occurred since only a minority of the bands from the genomic DNA had counterparts in the cosmid. Two hybridizing fragments from the cosmid, a 1.2 kb EcoRI fragment and a 1.0 kb PstI fragment, looked equal in size to hy-20 bridizing genomic fragments. The two fragments were sub-cloned from the cosmid and sequenced. Analysis of the sequence data showed that the fragments overlap. In total 1497 bp of sequence was obtained. Oligonucleotides encoding the Zn2+-finger were not contained within the sequence. A BamHI site was found close 25 to one end of the sequence in a region only covered by the EcoRI sub-clone, thus allowing the position of the sequenced genomic fragment relative to the Zn2+-finger to be determined. The primer 153468: CGGGATGAATTGTAGAGAGGC (SEQ ID NO: 41) was prepared by DNA Technology A/S (Denmark). The primer sequence 30 is contained within the 1497 bp fragment. It is found at the end closest to the ${\rm Zn}^{2+}\text{-finger}$ and points in that direction. Two primers both of prtT Zn2+-finger specific sequence and pointing either downstream (140358) or upstream (140359) were also prepared by DNA Technology A/S (Denmark). The sequence of

56

the two primers are as follows: 140358: CGCAAGCTTAAAACTCGGTGCGATC ID NO: (SEQ 42) · and CCTCGTGGATCAAGATCGCA (SEQ ID NO: 43). Two PCR reactions, one with the primers 153468 and 140358 and one with 153468 and 5 140359, respectively, were performed with genomic DNA as template. The reaction with 153468 and 140359 gave a band of approximately 1.1 kb, the other reaction gave no visible bands, when analysed on an agarose gel. The 1.1 kb fragment was cloned into pCR4Blunt-TOPO (Invitrogen) and sequenced. The fragment 10 contained part of the $\mathrm{Zn}^{2+}\text{-finger}$ and overlaps with the 1497bp fragment. Translation of the sequence showed that the region immediately upstream of the $\mathrm{Zn}^{2+}\text{-finger}$ encodes a polypeptide with homology to the N-terminal of prtT from A. niger.

15 Example 7

Isolation of the complete A. oryzae prtT gene

The remaining parts of the gene were cloned by two consecutive rounds of inverse PCR. In the first inverse PCR reaction the genomic DNA was restricted with EcoRV and re-ligated. 20 The PCR reaction was with the run primers 175653: GATGAAAAGAATAATCGGCGAG (SEQ ID NO: 44) and 175654: CGCGGCACACTACCCCCGTTG (SEQ ID NO: 45). The reaction resulted in the synthesis of a 1.9kb fragment, which was cloned into the pCR4Blunt-TOPO vector and sequenced. Analysis of the sequence 25 data showed that the fragment contains a gene with homology to the A. niger prtT gene and that the 3'end of the gene was missing. The second inverse PCR reaction was thus performed. primers were B0403G08: ATCTAGCTCAAGCATTAGCGGC (SEQ ID NO: 46) and B0403G09: AATTTCGGCCCTTTAGTGTCC (SEQ ID NO: 47). BglII re-30 stricted and re-ligated genomic DNA was used as template. A 2.4 kb fragment was obtained and cloned into the pCR4Blunt-TOPO vector and sequenced. Analysis of the sequence showed that the complete A. oyzae prtT gene had been obtained. The DNA sequence of the A. oryzae prtT gene is shown in SEQ ID NO: 48

57

and the deduced amino acid sequence of the encoded protein is shown in SEQ ID NO: 49.

Example 8

5 Disruption of the Aspergillus oryzae prtT gene.

The A. oryzae prtT gene was disrupted using a method of positive/negative selection. A disruption cassette consisting of 2kb of the A. oryzae prtT gene (SEQ ID NO: 48) with an insertion of the pyrG gene in the middle is cloned into a vector 10 (pDV8) containing the herpes simplex virus I tymidine kinase gene (HSV-tk) flanked by fungal expression signals. Expression of the tymidine kinase gene makes the host sensitive to 5flouro-2-deoxyuridine. A disrupted strain can be isolated by positive selection for the pyrG gene in a pyrG minus host and 15 deselection of the tymidine kinase gene on 5-flouro-2deoxyuridine. Since the tymidinie kinase gene and the pyrG gene are present in the same DNA fragment selection is for transformants in which a doubble cross-over event has happened. The system gives fewer transformants pr. Micro g of DNA than trans-20 formation with just a disruption cassette, but the frequency of transformants in which the desired homologous recombination event has occurred is much higher.

The pyrG gene used here is flanked by repeats enabling a later removal by selection for 5-fluoroorotic acid resistance. The pyrG gene is isolated from the plasmid pJaL554.

The pDV8 plasmid:

pDV8 was kindly provided by Matthew S. Sachs, University of Oregon, PO Box 91000, Portland, OR 97291-1000, USA. pDV8 30 (Fig. 7) is a pSP65 (Promega™) based plasmid containing the HSV-tk gene on a 1.2 kb BglII/BamHI fragment inserted between a 1.0 kb XhoI/BglII fragment of the A. nidulans gpd promoter and a 0.8 kb BamHI/HindIII fragment containing the A. nidulans trpC transcriptional terminator.

58

The A. nidulans gpd promoter and the trpC transcriptional terminator are taken from the plasmid pAN51-2 (Punt et al., (1990), Gene 93, p.101-109). The HSV-tk gene is described by McKnight S.L., (1980), Nucleic Acids Res. 8:5949-5964, Database 5 accession no. EMBL v00470, position 252-1479. The construction of pDV8 is described in Vaught-Alexander D (thesis) Expression of the herpes simplex virus type-1 thymidine kinase gene in Neurospora crassa, (1994), Oregon Graduate Institute of Science & Technology, University of Portland, PO Box 91000, Port-10 land, OR 97291-1000, USA. The sequence of pDV8 is included in this application as SEQ ID NO: 50. Single-, double- and multicopy A. oryzae transformants of pDV8 were isolated by transforming a pDV8 derivative containing the A. oryzae niaD gene into an A. oryzae niaD mutant. The copy number of the HSV-tk 15 gene was determined by Southern analysis. The transformants and the untransformed host were inoculated onto plates containing varying concentrations of 5-flouro-2'-deoxyuridine. From inspection of the growth on the plates it was decided to use 6 microM of 5-flouro-2'-deoxyuridine in the plates for future 20 positive/negative selection experiments. At this concentration none of the pDV8 transformants grew, while the untransformed host was only slightly inhibited.

Description of pJaL554:

25 PJaL554 was constructed by ligating the 316 bp Asp718-NheI fragment to the 5336 bp SpeI-SspBI fragment from the pyrG containing plasmid pSO2 (described in WO 97/35956). Thus, pJaL554 harbours the A. oryzae pyrG gene flanked by 316 bp repeats. The construction is illustrated in Fig. 8.

30

Construction of a prtT disruption plasmid in the pDV8 vector:

A PCR reaction is performed on chromosomal A. oryzae IFO4177 DNA with the primers B1042E05 and B1450E07. B1042E05: CGCGCGTATCCTATTGCC (SEQ ID NO: 51)

59

B1450E07: GCCGGAAATGTTGTACCTAC (SEQ ID NO: 52).

A fragment of 2078 basepairs is obtained and cloned into the pCR4Blunt-TOPO (Invitrogen™) vector. The resulting plasmid is sequenced with the standard M13 forward (-40) and reverse prim5 ers to ensure that the correct fragment is obtained. The PCR fragment is excised from the vector by the restriction enzyme EcoRV which cuts twice internally in the fragment. The cut sites are located at positions 1 and 1964 in SEQ ID NO: 48. The 1964 bp fragment is ligated with the pDV8 vector, which has been cut with HindIII and blunt ended by filling in the ends with the Klenow fragment of DNA polymerase I from E. coli and dNTP. The resulting plasmid is cut with HindIII, which is located in the prtT fragment (in the part encoding the Zn²+finger) at position 962 in SEQ ID NO: 48, dephosphorylated and ligated with the pyrG gene isolated from pJaL554 as a 2.5 kb HindIII fragment.

Selection of prtT disrupted strains:

The disruption plasmid described above was linearized with NotI and transformed into A. oryzae HowB101 (described in 20 WO 97/35956), a pyrG minus derivative of IFO4177. The transformation is done essential as described in EP 0 238 023. Transformants are selected on plates containing Coves salt solution (Cove DJ, (1966), Biochim. Biophys. Acta 113:51-56), 1 M sucrose for osmotic stabilization and as carbon source, 20g/L 25 agar, 10 mM NaNO3 and 6 microM 5-flouro-2-deoxyuridine (Sigma). The transformants are reisolated once on the same type of plates. Transformants carrying a disrupted prtT gene are identified by Southern blot analysis.

A strain carrying the prtT disruption is used as host for expression of a truncated PDI gene (Protein Disulfide isomerase gene) harbored on the expression plasmid pCaHj445 (described in US patent 5,879,664). pCaHj445 is transformed into the A. oryzae prtT disrupted strain by cotransformation with the plasmid p3SR2 containing the A. nidulans amdS gene. Transformation

60

and selection on acetamide plates is done essentially as described in EP 0 238 023. After reisolation the transformants are fermented in shake flasks or fermentors and the PDI protein is purified from the fermentation broth.

5

The invention described and claimed herein is not to be limited in scope by the specific embodiments herein disclosed, since these embodiments are intended as illustrations of several aspects of the invention. Any equivalent embodiments are intended to be within the scope of this invention. Indeed, various modifications of the invention in addition to those shown and described herein will become apparent to those skilled in the art from the foregoing description. Such modifications are also intended to fall within the scope of the appended claims.

Various references are cited herein, the disclosures of which are incorporated by reference in their entireties.

0-1	Form - PCT/RO/134 (EASY) Indications Relating to Deposited Microorganism(s) or Other Biologica	1
0-1-1	Material (PCT Rule 13bis) Prepared using	
0-1-1	Frepared using	PCT-EASY Version 2.91
		(updated 01.01.2001)
0-2	International Application No.	
		PCT/DK 01/00169
0-3	Applicant's or agent's file reference	10023-WO
1	The indications made below relate to the deposited microorganism(s) or other biological material referred to in the description on:	·
1-1	page	44
1-2	line	4-6
1-3	Identification of Deposit	14-0
1-3-1	Name of depositary institution	
	induction	DSMZ-Deutsche Sammlung von
1-3-2	Address of depositary institution	Mikroorganismen und Zellkulturen Gmb
	and the copositery monthly	Mascheroder Weg 1b, D-38124
1-3-3	Date of deposit	Braunschweig, Germany
1-3-4	Accession Number	14 July 1998 (14.07.1998)
1-4	Additional Indications	DSMZ 12294
1-5	Designated States for Which	NONE
	Indications are Made	all designated States
1-6	Separate Furnishing of Indications	NONE
	These indications will be submitted to the International Bureau later	
2	The indications made below relate to the deposited microorganism(s) or	
	other biological material referred to	
2-1	in the description on:	
2-2	line	44
2-3	Identification of Deposit	4-6
2-3-1		
-0-1	Name of depositary institution	DSMZ-Deutsche Sammlung von
2-3-2	Address of the transfer	Mikroorganismen und Zellkulturen GmbH
-0-2	Address of depositary Institution	Mascheroder Weg 1b, D-38124
	Bata of L	Braunschweig, Germany
2-3-3 2-3-4	Date of deposit	14 July 1994 (14.07.1994)
	Accession Number	DSMZ 12298
	Additional Indications	NONE
-5	Designated States for Which Indications are Made	all designated States
-6	Separate Furnishing of Indications	NONE

Claims:

10

15

20

- 1. An isolated nucleic acid sequence encoding a polypeptide having transcriptional activation activity, selected from the group consisting of:
 - (a) a nucleic acid sequence having at least 70% identity with the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48;
 - (b) a nucleic acid sequence encoding a polypeptide having an amino acid sequence which has at least 50% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49;
- a nucleic acid sequence which hybridizes under low (c) stringency conditions with (i) the nucleic acid sequence SEQ ID NO:1 or SEQ ID NO: 48, or (ii) complementary strand, wherein the low stringency conditions are defined by prehybridization hybridization at 42°C in 5x SSPE, 0.3% SDS, 200 g/ml sheared and denatured salmon sperm DNA, and 25% formamide, and wash conditions are defined by 50°C for 30 minutes in 2X SSC, 0.2% SDS;
 - (d) an allelic variant of (a), (b), or (c);
 - (e) a subsequence of (a), (b), (c), or (d), wherein the subsequence encodes a polypeptide fragment which has transcriptional activation activity; and
 - (f) a subsequence of (a), (b), (c), or (d), wherein the subsequence encodes a polypeptide with the amino acid sequence of SEQ ID NO:3.
- The nucleic acid sequence of claim 1 which has at least 70%, preferably at least 80%, more preferably at least 85%, even more preferably at least 90%, most preferably at least 95%, even most preferably at least 97%, and even more preferred

63

at least 99% identity with the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48.

- 3. The nucleic acid sequence of claim 1 which has the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48.
- The nucleic acid sequence of any of claims 1 to 3 which encodes a polypeptide comprising an amino acid sequence which has at least 50%, preferably at least 60%, preferably at least 70%, more preferably at least 80%, even more preferably at least 90%, and most preferably at least 95% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49.
- 15 5. The nucleic acid sequence of any of claims 1 to 4, wherein the nucleic acid sequence is obtained from a fungal cell or a yeast cell.
- 6. The nucleic acid sequence of claim 5, wherein the fungal cell is a filamentous fungal cell.
 - 7. The nucleic acid sequence of claim 6, wherein the filamentous fungal cell is an Aspergillus, Fusarium, Penicillium, or Trichoderma cell.

25

8. The nucleic acid sequence of claim 7, wherein the Aspergillus cell is a strain of Aspergillus niger or Aspergillus oryzae, or a respective synonym or teleomorph thereof.

30

9. The nucleic acid sequence of claim 8, wherein the Aspergillus cell is a strain of Aspergillus niger DSM 12298 or Aspergillus oryzae IFO4177.

64

- 10. The nucleic acid sequence of claim 7, wherein the Fusarium cell is a strain of Fusarium venenatum, or a synonym or teleomorph thereof.
- 5 11. The nucleic acid sequence of claim 5, wherein the yeast cell is a Hansenula, Pichia, or Saccharomyces cell.
 - 12. The nucleic acid sequence of any of claims 1 to 11 which encodes a polypeptide comprising the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49, or a fragment thereof, which has transcriptional activation activity.

- 13. The nucleic acid sequence of any of claims 1 to 11 which encodes a polypeptide comprising the amino acid sequence of SEQ ID NO:3.
- 14. The nucleic acid sequence of any of claims 1 to 13, which hybridizes under low, preferably medium, and more preferably high, stringency conditions to (i) the nucleic acid sequence set forth in SEQ ID NO:1 or SEQ ID NO: 48 or (ii) the respective complementary strand, or a subsequence thereof.
- 15. The nucleic acid sequence of claim 14, wherein low stringency conditions are defined by prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micro g/ml sheared and denatured salmon sperm DNA, and 25% formamide, and wash conditions are defined by 50°C for 30 minutes in 2X SSC, 0.2% SDS; medium stringency conditions are defined by prehybridization and hybridization at 42°C in 5x SSPE, 0.3% SDS, 200 microg/ml sheared and denatured salmon sperm DNA, and 35% formamide, and wash conditions are defined by 60°C for 30 minutes in 2X SSC, 0.2% SDS; and high stringency conditions are defined by prehybridization and hybridization at 42°C in 5x SSPE, 0.3% SDS, 200 microg/ml sheared and

denatured salmon sperm DNA, and 50% formamide, and wash conditions are defined by 70°C for 30 minutes in 2X SSC, 0.2% SDS.

- 5 16. The nucleic acid sequence of any of claims 1 to 15, which comprises the nucleic acid sequence encoding a polypeptide, which has transcriptional activation activity contained in the plasmid pEES harboured in *Escherichia coli* DSM 12294 or DNA sequence shown in SEQ ID NO: 48 encoding polypeptide shown in SEQ ID NO: 49.
 - 17. A nucleic acid construct comprising the nucleic acid sequence of any of claims 1 to 16 operably linked to one or more control sequences, which direct the production of the polypeptide in a suitable expression host.
 - 18. An expression vector comprising the nucleic acid construct of claim 17, a promoter, and transcriptional and translational stop signals.

20

30

- 19. A host cell comprising the nucleic acid construct of claim 17 or the expression vector of claim 18.
- 20. An isolated polypeptide selected from the group consisting of:
 - a polypeptide which is encoded in a nucleic acid (a) sequence which hybridizes under low stringency conditions with (i) the nucleic acid sequence of SEQ ID NO:1 or SEQ ID NO: 48; (ii) its complementary strand, or (iii) a subsequence of SEQ ID NO:1 or SEQ ID NO: 48 which encodes polypeptide fragment which has transcriptional activation activity, wherein the low stringency conditions are defined by prehybridization hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micro

g/ml sheared and denatured salmon sperm DNA, and 25% formamide, and wash conditions are defined at 50°C for 30 minutes in 2X SSC, 0.2% SDS;

- (b) a polypeptide having an amino acid sequence which has at least 50% identity with the amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 49;
- (c) an allelic variant of (a) or (b);
- (d) a fragment of (a), (b), or (c), wherein the fragment has transcriptional activation activity; and
- 10 (e) a polypeptide comprising the amino acid sequence of SEQ ID NO:3, or an allelic variant thereof.
- 21. The polypeptide of claim 20, comprising an amino acid sequence which has at least 50%, preferably at least 60%, preferably at least 70%, more preferably at least 80%, even more preferably at least 90%, more preferably at least 95%, even more preferred 97%, and most preferred 99% identity with the amino acid sequence of SEQ ID NO:2 or SEQ ID NO:49.

20

5

22. The polypeptide of claim 20 or 21, comprising the amino acid sequence of SEQ ID NO:2 or SEQ ID NO: 49, or a fragment thereof, wherein the fragment retains transcriptional activation activity.

25

23. The polypeptide of any of claims 20 to 22 which is encoded in the nucleic acid sequence contained in plasmid pEES which is contained in *Escherichia coli* DSM 12294 or the DNA sequence shown in SEQ ID NO: 48.

30

24. The polypeptide of any of claims 20 to 23 which comprises the amino acid sequence of SEQ ID NO:3.

25. A method for producing the polypeptide of any of claims 20 to 24 comprising (a) cultivating the host cell of claim 19 under conditions conducive for production of the polypeptide; and (b) recovering the polypeptide.

5

10

15

20

- 26. A fungal host cell useful for the production of a polypeptide, wherein the cell:
 - a) is a mutant of a parent fungal cell in which the parent cell comprises one or more DNA sequences encoding a protease, the transcription of which is activated by a transcriptional activator encoded by a nucleic acid sequence of any of claims 1 to 16; and
 - b) produces less of the transcriptional activator and the protease(s) than the parent cell when cultured under the same conditions.
- 27. The host cell of claim 26, wherein the reduced production of the transcriptional activator is obtained by modification or inactivation of a nucleic acid sequence present in the cell and necessary for expression of the transcriptional activator.
- 28. The host cell of claim 26 or 27, wherein the reduced production of the transcriptional activator is obtained by modification or inactivation of a control sequence required for the expression of the polypeptide.
 - 29. The host cell of claim 28, wherein the control sequence is a promoter sequence, or a functional part thereof.

30

30. The host cell of any of claims 26 to 29, wherein the nucleic acid sequence to be modified or inactivated is the sequence defined in any of claims 1 to 16.

68

PCT/DK01/00169

WO 01/68864

10

- 31. The host cell of any of claims 26 to 30, wherein the modification or inactivation is performed by specific or random mutagenesis, site-directed mutagensis, PCR generated mutagenesis, nucleotide insertion and/or substitution, gene interruption or gene replacement techniques, anti-sense techniques, or a combination thereof.
- 32. A fungal host cell useful for the production of a polypeptide, wherein the host cell is a mutant of a parent cell, in which the mutant:
 - a) produces more of a transcriptional activator encoded by a nucleic acid sequence of any of claims 1 to 16 than the parent cell when cultured under the same conditions, and
- b) comprises a DNA sequence encoding the polypeptide, the transcription of which is activated by the transcriptional activator.
- 33. The host cell of claim 32, wherein the host cell produces more of the transcriptional activator than the parent cell by introducing into the parent cell one or more copies of: (i) a nucleic acid sequence of any of claims 1 to 16, (ii) the nucleic acid construct of claim 17, or (iii) the expression vector of claim 18, whereby the host cell produces more of the polypeptide than the parent cell when cultured under the same conditions.
 - 34. The host cell of claim 32 or 33, wherein the nucleic acid acid sequence encoding the transcriptional activator is operably linked to a promoter, which is stronger than the corresponding promoter of the parent cell.
 - 35. The host cell of claim 34, wherein the promoter mediates the expression of a gene encoding an extracellular protease, preferably Aspergillus oryzae alkaline protease, A. oryzae

neutral metalloprotease, A. niger aspergillopepsin protease, Fusarium oxysporum trypsin-like protease or F. venenatum trypsin.

5 36. A fungal host cell useful for the production of a polypeptide, wherein the cell is a mutant of a parent cell in which the mutant comprises:

10

15

- a) a modification or inactivation of a transcriptional activator which is encoded in a native nucleic acid sequence of any of claims 1 to 16, or a regulatory sequence thereof, and
- b) (i) an inducible promoter operably linked to a nucleic acid sequence of any of claims 1 to 16, and (ii) a promoter sequence to which a transcriptional activator encoded by the nucleic acid sequence of any of claims 1 to 16 can bind, operably linked to a nucleic acid sequence encoding the polypeptide, wherein (i) and (ii) can be introduced simultaneously or sequentially.
- 20 37. The host cell of claim 36 wherein the native nucleic acid sequence, or a regulatory sequence thereof, is modified or inactivated by specific or random mutagenesis, site-directed mutagensis, PCR generated mutagenesis, nucleotide insertion and/or substitution, gene interruption or gene replacement techniques, anti-sense techniques, or a combination thereof.
 - 38. The host cell of claim 36 or 37, wherein the inducible promoter is selected from the group in which the induction is mediated by a carbon or nitrogen catabolite.
 - 39. The host cell of any of claims 33 to 38, which further comprises a promoter sequence, wherein the promoter sequence can be activated by the transcriptional activator and is

operably linked to the nucleic acid sequence encoding the polypeptide.

- 40. The host cell of any of claims 33 to 39, wherein the promoter sequence, or a functional part thereof, is from a protease gene.
- 41. The host cell of any of claims 33 to 40, wherein the protease gene is Fusarium oxysporum trypsin-like protease gene, Aspegillus oryzae alkaline protease gene, Aspergillus niger pacA gene, Aspergillus oryzae alkaline protease gene, A. oryzae neutral metalloprotease gene, A. niger aspergillopepsin protease gene, or F. venenatum trypsin gene.

- 42. The host cell of any of claims 26 to 41, wherein the host cell comprises at least one copy of a nucleic acid sequence encoding the polypeptide.
- 20 43. The host cell of any of claims 26 to 42, wherein the host cell produces less of a native protease or a combination of native proteases than the parent cell when cultured under identical conditions.
- 25 44. The host cell of any of claims 26 to 43, wherein the activity of the protease is assayed by the degradation of ³H-labelled sperm whale myoglobin at pH 4.
 - 45. A method of producing a polypeptide, comprising:
- (a) cultivating the host cell of any of claims 26 to 44, wherein the host cell harbours a gene encoding the polypeptide, in a nutrient medium suitable for production of the polypeptide; and

71

- (b) recovering the polypeptide from the nutrient medium of the mutant cell.
- 46. The method of claim 45, wherein the polypeptide is native to the parent cell.
 - 47. The method of claim 45, wherein the polypeptide is heterologous to the parent cell.
- 10 48. The method of claim 45, wherein the polypeptide is an antibody or portions thereof, an antigen, a clotting factor, an enzyme, a hormone or a hormone variant, a receptor or portions thereof, a regulatory protein, a structural protein, a reporter, or a transport protein.

15

- 49. The method of claim 48, wherein the enzyme is an oxidoreductase, transferase, hydrolase, lyase, isomerase, or ligase.
- 20 50. The method of claim 49, wherein the enzyme aminopeptidase, amylase, carbohydrase, carboxypeptidase, catalase, cellulase, chitinase, cutinase, deoxyribonuclease, dextranase, esterase, alpha-galactosidase, betagalactosidase, glucoamylase, alpha-glucosidase, beta-25 glucosidase, haloperoxidase, invertase, laccase, lipase, mannosidase, mutanase, oxidase, pectinolytic enzyme,

ribonuclease, transglutaminase, or xylanase.

peroxidase, phytase, polyphenoloxidase, proteolytic enzyme,

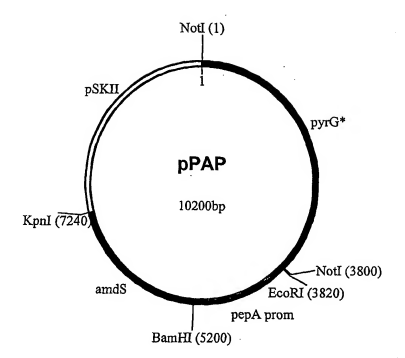


Fig. 1

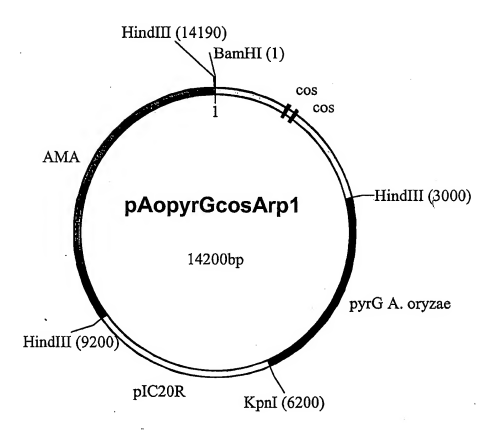


Fig. 2

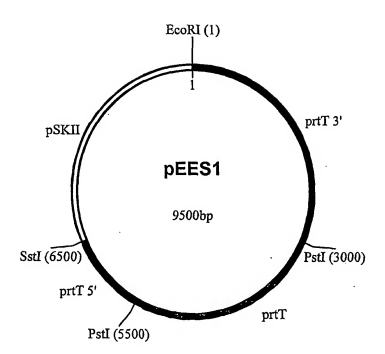


Fig. 3

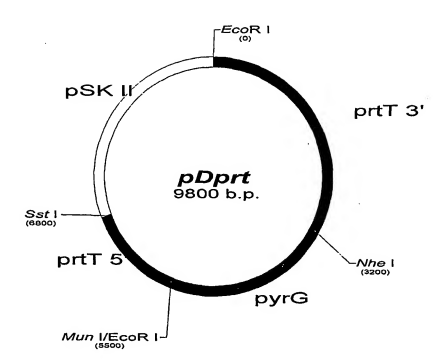


Fig. 4

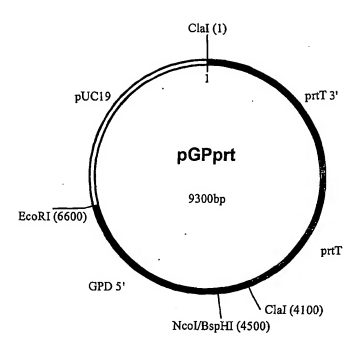


Fig. 5

WO 01/68864

6/8

ICA217 ATGACCGCTT GTCATACCTG CCGCAAGCTT AAAACTCGGT ICA218 ATGACTGCTT GCCACACCTG CCGCAAGCTT AAAACTCGGT

ICA217 GCGATCTTGA TCCACGAGGG CATGCCTGCC GCCGCTGCCT (SEQ ID NO: 53)
ICA218 GCGATCTTGA TCCACGAGGG CATGCCTGCC GCCGCTGCCT (SEQ ID NO: 54)

Fig. 6

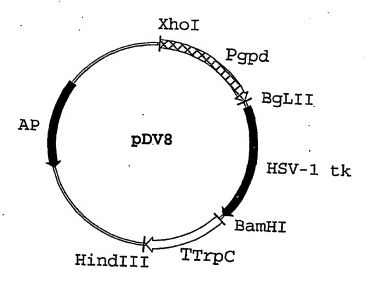


Fig. 7

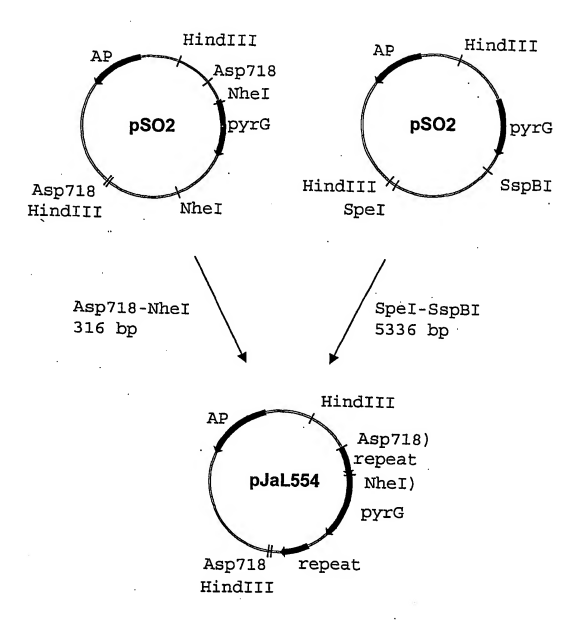


Fig. 8

SEQUENCE LISTING

```
<110> Novo Nordisk A/S
     <120>
 5
     <130>
     <160> 54
10
     <170> PatentIn Ver. 2.1
     <210> 1
     <211> 4098
     <212> DNA
15
     <213> E. coli DSM 12294
     <220>
     <221> exon
     <222> (977)..(1204)
20
     <220>
     <221> exon
     <222> (1317)..(1718)
25
    <220>
     <221> exon
     <222> (1777)..(2202)
    <220>
30
     <221> exon
     <222> (2253)..(3116)
     <220>
     <221> exon
     <222> (3170)..(3247)
35
     <400> 1
     ttggtgctgg aaagcccatt taagggatct tataaggtaa ttgccaatgt tcagtcgcct 60
40
    atggtctttg tcgagagaaa ctctttctcg ttaagatcta catgatcgct tttgattttc 120
     tetgggttea egeggtaett teteceegte aateceeaac egetgttgtg cetgaceate 180
     aatgtggaac ggataagggg acaagagaaa ttgaaggagc gatcataaaa agctaatttt 240
45
    ggtttattat ttttttttt tataaaactc aaaaaagaaa acgaaaacga aaaaggaaaa 300
     aagaaaaggt aaaatggaaa aagaaaggcg gtcatcactt ccaataacca tcagccaaag 360
50
    atacagacga gttactgacc ttcttatcct ggacttccgc ccgatccata tcttcatgat 420
     aagcagggaa ccgaacaaat caacgccaac ttcagcggca gttcctcact aatttcccac 480
     ttcccacegg cgtcattttg gtcccaaccc cctccctgga agcageggga tttagttacg 540
55
    atceggttta categgagae teggaaaata ecatagegea tgecaateaa aacceeteee 600
    agggtgactg gccagtatca cgacccattg tttctatctt tctagaagac ctgcagggac 660
60
    atggattggc tggccgccgt gctgccgtcc attagcgtct accccaggtc aagaacggac 720
```

	tggac	ggacc	cataa	accaa	at c	taac	caaag	gcc	aatti	tcgt	caa	ttcc	cag	ctgg	cgagca	780
5	caato	ccatt	cccag	gggtl	tg g	ccgc	caact	gti	taaa	aggc	act	atgt	gtc	tete	cacctg	840
	cccgc	cccc	tcgat	ggc	ct g	cgcgt	taata	a act	tatte	ctac	tgc	tttt	tgc (ctct	tacttg	900
	cctca	ttatt	agtat	ttta	ac to	ctact	ctcc	aga	attgo	cctg	cca	gcaat	ttg q	gtcc	aaagtg	960
10	gactt	tgttt	gatga	ac at	tg a	ct c	ga ac	cc gt	tg ga	ac ga	ag at	tc a	aa ta	ac ga	aa acg	1012
	cct t	ct tca	tgg	gag	cac	aag	agc	ttg	gac	gtt	gcc	gag	gat	ggc	agg	1060
15	cga c	ta gct	ccc	cat	tcc	gac	act	gct	cgt	ccg	aaa	ggc	cgc	ata	cga	1108
13	cga t	cg atg	act	gcc	tgt	cac	aca	tgt	cgg	aag	ctt	aaa	act	aga	tgt	1156
	gat c	ta gat	ccg	cgc	ggt	cat	gcg	tgc	cgt	cgc	tgt	cta	tct	cta	agg	1204
20	tcaga	ggcac	tacct	acct	g c	cagt	gaag	g ctt	tgt	cctt	ctg	aacg	cga (catg	atacta	1264
	gtcgt	ggaat :	ataad	ctgto	CC C	aact	tgct	gad	cagto	ccac	aata	atct	tta 🤉	ga at	tc gat	1322
25	tgt a	ag ctg	cct	gaa	acg	acc	gac	cgc	ttc	caa	gac	agt	gct	gcg	atg	1370
	tgg c	ca gac	gcc	acc	tcg	gca	att	ccc	tcc	atc	gag	gag	cgc	ctc	acc	1418
	tcc c	ta gaa	aga	tgc	atg	agg	gag	atgi	acg	ggc	atg	atg	cga	cag	atg	1466
30	cta g	at cac	tcc	cca	ggt	ttc	gca	aat	gcc	tcg	gtt	ccg	cat	ttg	acc	1514
	aaa a	gc atc	atc	acg	gat	gaa	acc	gcc	tcg	atg	gag	gga	agc	ccg	tcg	1562
35	tcc c	cc ttc	ctg	cct	aag	ccc	gtt	cgc	ctc	att	cag	gac	ctc	cag	tcc	1610
	gac t	tc ttc	gga	gaa	gca	gag	act	tcc	ccc	gtt	gac	tcc	cct	ctc	tcc	1658
	agc g	at ggt	aac	gcc	aag	ggc	gct	atc	gac	tct	aag	cta	tcc	ctc	aaa	1706
40	ttg t	tg caa	acg	tate	gggta	ata d	ctga	attga	ac aa	atta	ccaa	a aaq	gctg	ctaa		1758
	tcctt	ggcgc a	aaato	cagg	ttt	gtc	gat	cac	ttt	ggc	gct	tgc	gtt	tcc	att	1809
45	tac a	at ctc	tcc	gac	atc	cac	aac	gac	atg	aaa	gcc	ccc	gac	tct	tta	1857
	ctg t	at aat	act	gca	tgc	ctt	cta	gct	tca	cgc	tat	gta	ccg	9 99	ata	1905
	ccg a	ca tct	acc	gtg	cat	gct	ata	tac	ctt	caa	gtg	cga	cat	gca	gta	1953
50	gtc a	at att	ttg	tgg	gaa	aaa	cca	ccc	ctg	aag	tat	gag	acc	ctc	caa	2001
	gca c	tt gca	ctt	ctc	tgt	ctc	tgg	cca	gca	acc	gcc	cag	aaa	gag	cca	2049
55	ccc a	tg gac	aġc	tgg	ctg	ctg	agt	ggt	atc	tca	att	aac	cat	gca	att	2097
	atc g	eg ete	gat	ttc	cta	aac	tat	gcg	ccc	tcg	gaa	gtc	atg	gtg	gac	2145
	aat g	aa acg	gct	gcg	cag	ctg	cgg	cta	tgg	aat	aca	tat	tgc	ttg	aca	2193
60	cag c	ta cag	tggg	gtttc	at o	ctaac	atct	c co	gtco	agaa	a gat	agct	aac			2242

	aago	ettta	agt t	tt g	geg g	gtc 9	399 a	aat g	geg d	egt d	cct 1	ttc (cat a	atc .	cag (caa	2291
															gca		2339
5															atg		2387
															ctt		2435
															ttc		2483
10	aag	aag	cct	gtt	ctt	gtt	tcc	cgg	gga	cta	cca	ctg	acg	aga	gca	aca	2531
	gct	999	gaa	agt	tcc	aca	ttg	gag	ctg	agc	ctt	tgg	ttc	tgc	cag	aca	2579
15	ctc	ctt	cac	cgc	aca	gca	atg	agg	ctt	cag	CCC	aga	tcc	gac	agg	ctc	2627
	gca	tct	gag	gtt	ctg	caa	acc	tca	cgt	ctg	ata	ata	tcg	cgg	ttc	ctc	2675
	cag	atc	cgg	tac	tct	acc	gca	tta	agc	ctt	gtc	gac	caa	gtc	tat	ttc	2723
20	att	gtc	ggc	tac	gct	gca	ctg	aat	ctg	tgc	gat	ttc	aat	ctt	atg	gac	2771
	ccg	ctt	atc	gag	caa	gtg	cag	atg	ttc	ctg	ctg	cat	ctc	tcc	ccg	aac	2819
25	gaa	gac	cac	atc	gcc	tac	cgg	ttt	tcg	tgc	atg	gtc	gcc	gag	ttc	aag	2867
	cgg	cga	tgt	ggc	agt	gcg	gaa	tgc	aat	gac	cca	tca.	tcc	act	gtc	aag	2915
20	ggg	tct	ccg	tta	tca	tcc	tac	ggc	gac	agt	cgt	aag	atg	agc	atg	999	2963
30	caa	gca	ccg	ttc	atg	cca	ccg	ctc	atg	gat	ggc	atg	atc	gag	ggg	tac	3011
	ggc	ttc	gag	caa	ctg	atg	cca	gaa	gtc	atg	ccg	agt	tcc	ttt	ccg	gat	3059
35	999	ata	ctc	aac	gga	atg	cct	gtg	act	aaa	cta	gca	gcg	tat	cgg	tca	3107
	gcg	acg	ctg	taag	gtaat	.cg a	gato	gggt	it go	gaaag	ggaca	a tga	agtg	3999			3156
40	tggt	ggtg	ggt a	igt a	igc a	igt a	ac a	acc a	agg g	gat c	gat a	aac o	etg d	cag (cgg t	gg	3205
40	ttt	agt	tcc	tgc	cca	tgg	gct	gaa	cta	aaa	ccc	cga	acc	tag			3247
	cato	gatga	acg t	gcaa	cgaa	a gg	jatca	ataac	caa	aggco	caag	taaa	atact	caa a	aataa	aataa	3307
45	tata	atto	ca c	cacga	tcca	c ta	ccac	ccacc	acc	acco	ggat	ccat	cago	gtt q	gcctt	cctgc	3367
	acag	gcct	at t	tagt	taga	a as	lccc	gtgcc	acc	jaaac	atc	acgt	aatt	ga g	geget	tttgc	3427
50	ttcc	ttgc	aa c	ttaa	acaa	c cc	cata	agaca	cto	ctcac	att	caca	atgco	caa a	actac	ctaact	3487
50	ccta	ctga	ecc a	ccag	ctgo	a gg	aago	cago	cag	gccac	cat	ttc	ctaat	cg g	gatat	atctc	3547
	cgaa	acgt	ac g	jcttt	cctc	c tt	tgtt	cgga	ccc	gttcc	gtg	ccto	cgcg	gga g	gagtt	gaacg	3607
55	agto	agaa	ca c	atto	tttt	c gt	ttct	atcg	, ttt	cttt	tcc	aagg	gcago	cag a	agaga	acgaac	3667
	aagt	cagt	gc t	tgct	aact	a ac	ttac	ccct	cag	gcatt	tta	gtaa	acta	act a	attta	aggaaa	3727
60	gagt	aato	at t	cato	gaag	a ca	agat	gttt	att	tete	cga	tcga	eccaa	aac a	aaaa	acgttc	3787
	aggt	agac	ta a	ıgtag	tagt	a gt	agta	tgto	ttt	gaco	cct	ttac	etcca	act a	atccc	gttgac	3847

	-50.	-cuc	age .	ag ca	ag cu	AC C	accu	aacca	ı gcı	-gcc;	Jayy	aya	yaa	agu '	gagu	399 1 99	3907
5	gag	ccgg	agg a	atgc	cgcc	ga g	aatt	attaa	a gto	gato	catt	gct	agtta	agt	tatc	ttttca	3967
	tgai	tgag	gag a	agga	agga	ga g	9999	gacgg	g gat	taga	agaa	ata	aacti	ttt	ctct	ccaatt	4027
	aati	tatc	tgg ;	atta	attaa	aa a	cttg	gagag	g gag	gggta	aggg	gag	ttggg	gta	ttggi	tatgtt	4087
10	gct	gtga	atg 1	t													4098
15	<21:	0> 2 1> 60 2> PI 3> E	66 RT	i DSI	M 12:	294											
20		3> De	escr	iptio	on of	E Ar	tifi	cial	Seqı	ience	e: MI	3 L12 :	13				
25			Arg	Thr	Val 5	Asp	Glu	Ile	Lys	Tyr 10	Glu	Thr	Pro	Ser	Ser 15	Trp	
	Glu	His	Lys	Ser 20	Leu	Asp	Val	Ala	Glu 25	Asp	Gly	Arg	Arg	Leu 30	Ala	Pro	
30	His	Ser	Asp 35	Thr	Ala	Arg	Pro	Lys 40	Gly	Arg	Ile	Arg	Arg 45	Ser	Met	Thr	
	Ala	Cys 50	His	Thr	Cys	Arg	Lys 55	Leu	Lys	Thr	Arg	Cys 60	Asp	Leu	Asp	Pro	
35	65					70					75				Cys	80	
40					85					90					Trp 95		
				100					105					110	Ser		
45			115					120					125		Leu	_	
		130					135					140			Lys		
50	Ile 145	Ile	Thr	Asp	Glu	Thr 150	Ala	Ser	Met	Glu	Gly 155	Ser	Pro	Ser	Ser	Pro 160	
55					165					170					Asp 175		
				180					185	_				190	Ser	_	
60	Gly	Asn	Ala 195	ГÀЗ	Gly	Ala	Ile	Asp 200	Ser	Lys	Leu	Ser	Leu 205	Lys	Leu	Leu	

	GIII	210	FIIE	val	Авр	птр	215	GIY	Ala	Сув	vai	220	TTE	ıyr	Asn	Let
5	Ser 225	Asp	Ile	His	Asn	Asp 230	Met	Lys	Ala	Pro	Asp 235	Ser	Leu	Leu	Tyr	Asi 240
	Thr	Ala	Cys	Leu	Leu 245	Ala	Ser	Arg	Tyr	Val 250	Pro	Gly	Ile	Pro	Thr 255	Sei
10	Thr	Val	His	Ala 260	Ile	Tyr	Leu	Gln	Val 265	Arg	His	Ala	Val	Val 270	Asn	Ile
15	Leu	Trp	Glu 275	Lys	Pro	Pro	Leu	Lys 280	Tyr	Glu	Thr	Leu	Gln 285	Ala	Leu	Ala
13	Leu	Leu 290	Cys	Leu	Trp	Pro	Ala 295	Thr	Ala	Gln	Lys	Glu 300	Pro	Pro	Met	Ası
20	Ser 305	Trp	Leu	Leu	Ser	Gly 310	Ile	Ser	Ile	Asn	His 315	Ala	Ile	Ile	Ala	Let 320
	Asp	Phe	Leu	Asn	Tyr 325	Ala	Pro	Ser	Glu	Val 330	Met	Val	Asp	Asn	Glu 335	Thi
25	Ala	Ala	Gln	Leu 340	Arg	Leu	Trp	Asn	Thr 345	Tyr	Сув	Leu	Thr	Gln 350	Leu	His
30	Phe	Ala	Val 355	Gly	Asn	Ala	Arg	Pro 360	Phe	His	Ile	Gln	Gln 365	Arg	Tyr	Leu
	Asp	His 370	Сув	Pro	Arg	Ile	Leu 375	Glu	His	Pro	Ala	Ala 380	Thr	Leu	Glu	Asp
35	Ala 385	Arg	Val	Val	Ala	Glu 390		Gln	Leu	тут	Leu 395	Met	Thr	Leu	Arg	Let 400
	Gln	Ser	Asn	Ser	Ser 405	Arg	Met	Arg	Leu	Ala 410	Asp	Leu	Asp	Tyr	Glu 415	Glu
40	Ile	Glu	Arg	Trp 420	Lys	Arg	Glu	Trp	Ala 425	His	Leu	Phe	Cys	Lys 430	Lys	Pro
45	Val	Leu	Val 435	Ser	Arg	Gly	Leu	Pro 440	Leu	Thr	Arg	Ala	Thr 445	Ala	Gly	Glu
43	Ser	Ser 450	Thr	Leu	Glu	Leu	Ser 455	Leu	Trp	Phe	Сув	Gln 460	Thr	Leu	Leu	His
50	Arg 465	Thr	Ala	Met	Arg	Leu 470	Gln	Pro	Arg	Ser	Asp 475	Arg	Leu	Ala	Ser	Glu 480
	Val	Leu	Gln	Thr	Ser 485	Arg	Leu	Ile	Ile	Ser 490	Arg	Phe	Leu	Gln	Ile 495	Arg
55	Tyr	Ser	Thr	Ala 500	Leu	Ser	Leu	Val	Asp 505	Gln	Val	Tyr	Phe	Ile 510	Val	Gly
•	Tyr	Ala	Ala 515	Leu	Asn	Leu	Cys	Asp 520	Phe	Asn	Leu	Met	Asp 525	Pro	Leu	Ile
60	Glu	Gln	Val	Gln	Met	Phe	Leu	Leu	His	Leu	Ser	Pro	Asn	Glu	geA	His

		530					535					540			•		
5	Ile 545	Ala	Tyr	Arg	Phe	Ser 550	Сув	Met	Val	Ala	Glu 555	Phe	Lys	Arg	Arg	Сув 560	
J	Gly	Ser	Ala	Glu	Cys 565	Asn	Asp	Pro	Ser	Ser 570	Thr	Val	Lys	Gly	Ser 575	Pro	
10	Leu	Ser	Ser	Tyr 580	Gly	Asp	Ser	Arg	Lys 585	Met	Ser	Met	Gly	Gln 590	Ala	Pro	
	Phe	Met	Pro 595	Pro	Leu	Met	Asp	Gly 600	Met	Ile	Glu	Gly	Tyr 605	Gly	Phe	Glu	
15	Gln	Leu 610	Met	Pro	Glu	Val	Met 615	Pro	Ser	Ser	Phe	Pro 620	Asp	Gly	Ile	Leu	
20	Asn 625	Gly	Met	Pro	Val	Thr 630	Gly	Leu	Ala	Ala	Tyr 635	Arg	Ser	Ala	Thr	Leu 640	
	Ser	Ser	Asn	Thr	Arg 645	Asp	Asp	Asn	Leu	Gln 650	Arg	Trp	Phe	Ser	Ser 655	Сув	
25	Pro	Trp	Ala	Glu 660	Leu	Lys	Pro	Arg	Thr 665	Pro							
30	<212)> 3 L> 35 2> PF 3> As	$\mathbf{T}^{\mathbf{S}}$	g ill ı	ıs ni	ger											
35	<220 <223		escri	ptic	n of	Art	ific	cial	Sequ	ience	e: Pe	epApı	c				
40	<400 Met 1		Ala	Cys	His 5	Thr	Cys	Arg	Lys	Leu 10	Lys	Thr	Arg	Сув	Asp 15	Leu	
	Asp	Pro	Arg	Gly 20	His	Ala	Сув	Arg	Arg 25	Сув	Leu	Ser	Leu	Arg 30	Ile	Asp	
45	Сув	ГУв	Leu 35														
50	<212	> 25 > DN	IA	.cial	. Sec	quenc	e										
55	<220 <223	> De	scri	ptic alle	n of	Art	ific	ial	Sequ	ience	e: PC	CR F1	cagme	ent			
50	gact	itgga :ggac	:gg a	ICCC8	taac	c aa	itcta	acca	ı aaç	gccaa	ittt	cgto	caatt	cc c	agct	igaacg :ggcga :tccac	120

```
ctgcccgccc ccctcgatgg cctgcgcgta ataactattc tactgctttt tgcctcttac 240
     ttgcctcatt attagtattt tactctactc tccagattgc ctgccagcaa ttggtccaaa 300
     gtggactttg tttgatgaca tgactcgaac cgtggacgag atcaaatacg aaacgccttc 360
     ttcatgggag cacaagagct tggacgttgc cgaggatggc aggcgactag ctccccattc 420
     cgacactgct cgtccgaaag gccgcatacg acgatcgatg actgcctgtc acacatgtcg 480
     gaagettaaa actagatgtg atctagatee gegeggteat gegtgeegte getgtetate 540
     tctaaggtca gaggcactac ctacctgcca gttgaagctt tgtccttctg aacgcgacat 600
     gatactagtc gtggaatata actgtcccaa ctttgctgac agtccacaat atctttagaa 660
     tegattgtaa getgeetgaa acgacegace getteeaaga eagtgetgeg atgtggeeag 720
     acgccacctc ggcaattccc tccatcgagg agegcctcac ctccctagaa agatgcatga 780 gggagatgac gggcatgatg cgacagatgc tagatcactc cccaggtttc gcaaatgcct 840
     cggttccgca tttgaccaaa agcatcatca cggatgaaac cgcctcgatg gagggaagcc 900
     egtegteece etteetgeet aagecegtte geeteattea ggaeeteeag teegaettet 960
     teggagaage agagaettee eeegttgaet eeeetetete cagegatggt aacgeeaagg 1020
15
     gegetatega etetaageta teeeteaaat tgttgeaaac gtatgggtat acetgattga 1080
     caattaccaa aaagetgeta ateettggeg caaatcaggt ttgtegatea etttggeget 1140
     tgcgtttcca tttacaatct ctccgacatc cacaacgaca tgaaagcccc cgactcttta 1200
     ctgtataata ctgcatgcct tctagcttca cgctatgtac cggggatacc gacatctacc 1260
     gtgcatgcta tataccttca agtgcgacat gcagtagtca atattttgtg ggaaaaacca 1320
20
     cccctgaagt atgagaccct ccaagcactt gcacttctct gtctctggcc agcaaccgcc 1380
     cagaaagagc cacccatgga cagctggctg ctgagtggta tctcaattaa ccatgcaatt 1440
     ategegeteg atttectaaa etatgegeee teggaagtea tggtggacaa tgaaaegget 1500
     gcgcagctgc ggctatggaa tacatattgc ttgacacagc tacagtgggt ttcatctaag 1560
     atctcccgtc cagaagatag ctaacaagct ttagttttgc ggtcgggaat gcgcgtcctt 1620
25
     tccatatcca gcaaagatac cttgaccact gcccacggat actggagcac ccagcagcaa 1680
     ctctggagga cgcaagggtt gtagcagaaa tacagttgta tttgatgaca ttgcggctcc 1740
     agaggagtg ggctcacctt ttctgtaaga agcctgttct tgtttcccgg ggactaccac 1860
     tgacgagagc aacagctggg gaaagttcca cattggagct gagcctttgg ttctgccaga 1920
30
     cacteettea eegeacagea atgaggette ageceagate egacaggete geatetgagg 1980
     ttctgcaaac ctcacgtctg ataatatcgc ggttcctcca gatccggtac tctaccgcat 2040
     taagcettgt cgaccaagte tattteattg teggetacge tgcactgaat etgtgegatt 2100
     tcaatcttat ggacccgctt atcgagcaag tgcagatgtt cctgctgcat ctctccccqa 2160
     acgaagacca categoetae eggttttegt geatggtege egagtteaag eggegatgtg 2220
35
     gcagtgcgga atgcaatgac ccatcatcca ctgtcaaggg gtctccgtta tcatcctacg 2280
     gcgacagtcg taagatgagc atggggcaag caccgttcat gccaccgctc atggatggca 2340
     tgatcgaggg gtacggcttc gagcaactga tgccagaagt catgccgagt tcctttccgg 2400
     atgggatact caacggaatg cctgtgactg ggctagcagc gtatcggtca gcgacgctgt 2460
     aagtaatega gategggttg gaaaggaeat gagtgggggt ggtggtggta gtageagtaa 2520
40
     caccagggat gataacctgc ag
                                                                         2542
     <210> 5
     <211> 665
     <212> PRT
     <213> Aspergillus niger
     <400> 5
     Met Thr Arg Thr Val Asp Glu Ile Lys Tyr Glu Thr Pro Ser Ser Trp
50
     Glu His Lys Ser Leu Asp Val Ala Glu Asp Gly Arg Arg Leu Ala Pro
     His Ser Asp Thr Ala Arg Pro Lys Gly Arg Ile Arg Arg Ser Met Thr
     Ala Cys His Thr Cys Arg Lys Leu Lys Thr Arg Cys Asp Leu Asp Pro
60
```

Arg Gly His Ala Cys Arg Arg Cys Leu Ser Leu Arg Ile Asp Cys Lys

	65					70					75					80
5	Leu	Pro	Glu	Thr	Thr 85	Asp	Arg	Phe	Gln	Asp 90	Ser	Ala	Ala	Met	Trp 95	Pro
J	Asp	Ala	Thr	Ser 100	Ala	Ile	Pro	Ser	Ile 105	Glu	Glu	Arg	Leu	Thr 110	Ser	Pro
10	Glu	Arg	Сув 115	Met	Arg	Glu	Met	Thr 120	Gly	Met	Met	Arg	Gln 125	Met	Leu	Asp
	His	Ser 130	Pro	Gly	Phe	Ala	Asn 135	Ala	Ser	Val	Pro	His 140	Leu	Thr	Гуз	Ser
15	Ile 145	Ile	Thr	Asp	Glu	Thr 150	Ala	Ser	Met	Glu	Gly 155	Ser	Pro	Ser	Ser	Pro 160
20	Phe	Leu	Pro	Lys	Pro 165	Val	Arg	Leu	Ile	Gln 170	qaA	Leu	Gln	Ser	Asp 175	Phe
	Phe	Gly	Glu	Ala 180	Glu	Thr	Ser	Pro	Val 185	Asp	Ser	Pro	Leu	Ser 190	Ser	Asp
25	Gly	Asn	Ala 195	Lys	Gly	Ala	Ile	Asp 200	Ser	Lys	Leu	Ser	Leu 205	Lys	Leu	Leu
	Gln	Thr 210	Phe	Val	Asp	His	Phe 215	Gly	Ala	Cys	Val	Ser 220	Ile	Tyr	Asn	Leu
30	Ser 225	Asp	Ile	His	Asn	Asp 230	Met	Lys	Ala	Pro	Asp 235	Ser	Leu	Leu	Tyr	Asn 240
35	Thr	Ala	Cys	Leu	Leu 245	Ala	Ser	Arg	Tyr	Val 250	Pro	Gly	Ile	Pro	Thr 255	Ser
	Thr	Val	His	Ala 260	Ile	Tyr	Leu	Gln	Val 265	Arg	His	Ala	Val	Val 270	Asn	Ile
40	Leu	Trp	Glu 275	Lys	Pro	Pro	Leu	Lys 280	Tyr	Glu	Thr	Leu	Gln 285	Ala	Leu	Ala
	Leu	Leu 290	Cys	Leu	Trp	Pro	Ala 295	Thr	Ala	Gln		Glu 300	Pro	Pro	Met	Asp
45	Ser 305	Trp	Leu	Leu	Ser	Gly 310	Ile	Ser	Ile	Asn	His 315	Ala	Ile	Ile	Ala	Leu 320
50	Asp	Phe	Leu	Asn	Tyr 325	Ala	Pro	Ser	Glu	Val 330	Met	Val	Asp	Asn	Glu 335	Thr
	Ala	Ala	Gln	Leu 340	Arg	Leu	Trp	Asn	Thr 345	Tyr	Cys	Leu	Thr	Gln 350	Leu	His
55	Phe	Ala	Val 355	Gly	Asn	Ala	Arg	Pro 360	Phe	His	Ile	Gl'n	Gln 365	Arg	Tyr	Leu
	Asp	His 370	Сув	Pro	Arg	Ile	Leu 375	Glu	His	Pro		Ala .380	Thr	Leu	Glu	Asp
60	Ala 385	Arg	Val	Val	Ala	Glu 390	Ile	Gln	Leu	Tyr	Leu	Met	Thr	Leu	Arg	Leu

	GIn	Ser	Asn	Ser	Ser 405	Arg	Met	Arg	Leu	Ala 410	Asp	Leu	Asp	Tyr	Glu 415	Glu
5	Ile	Glu	Arg	Trp 420	Lys	Arg	Glu	Trp	Ala 425	His	Leu	Phe	Сув	Lys 430	Lys	Pro
10	Val	Leu	Val 435	Ser	Arg	Gly	Leu	Pro 440	Leu	Thr	Arg	Ala	Thr 445	Ala	Gly	Glu
	Ser	Ser 450	Thr	Leu	Glu	Leu	Ser 455	Leu	Trp	Phe	Cys	Gln 460	Thr	Leu	Leu	His
15	Arg 465	Thr	Ala	Met	Arg	Leu 470	Gln	Pro	Arg	Ser	Asp 475	Arg	Leu	Ala	Ser	Glu 480
	Val	Leu	Gln	Thr	Ser 485	Arg	Leu	Ile	Ile	Ser 490	Arg	Phe	Leu	Gln	Ile 495	Arg
20	Tyr	Ser	Thr	Ala 500	Leu	Ser	Leu	Val	Asp 505	Gln	Val	Tyr	Phe	Ile 510	Val	Gly
25	Tyr	Ala	Ala 515	Leu	Asn	Leu	Cys	Asp 520	Phe	Asn	Leu	Met	Asp 525	Pro	Leu	Ile
	Glu	Gln 530	Val	Gln	Met	Phe	Leu 535	Leu	His	Leu	Ser	Pro 540	Asn	Glu	Asp _.	His
30	Ile 545	Ala	Tyr	Arg	Phe	Ser 550	Сув	Met	Val	Ala	Glu 555	Phe	ГÀЗ	Arg	Arg	Сув 560
	Gly	Ser	Ala	Glu	Сув 565	Asn	Asp	Pro	Ser	Ser 570	Thr	Val	ГЛя	Gly	Ser 575	Pro
35	Leu	Ser	Ser	Tyr 580	Gly	Asp	Ser	Arg	Lys 585	Met	Ser	Met	Gly	Gln 590	Ala	Pro
40	Phe	Met	Pro 595	Pro	Leu	Met	Asp	Gly 600	Met	Ile	Glu	Gly	Tyr 605	Gly	Phe	Glu
	Gln	Leu 610	Met	Pro	Glu	Val	Met 615	Pro	Ser	Ser	Phe	Pro 620	_	Gly	Ile	Leu
45	Asn 625	Gly	Met	Pro	Val	Thr 630	Gly	Leu	Ala	Ala	Tyr 635	Arg	Ser	Ala	Thr	Leu 640
	Ser	Ser	Asn	Thr	Arg 645	Asp	Asp	Asn	Leu	Gln 650	Arg	Trp	Phe	Ser	Ser 655	Cys
50	Pro	Trp	Ala	Glu 660	Leu	Lys	Pro	Arg	Thr 665							
55	-210)														

60 <220>
 <223> Description of Artificial Sequence: PepApr

	cggaattcgc atgctggagg tgcttctaa	29
5		
	<210> 7	
	<211> 33	
	<212> DNA <213> Artificial Sequence	
10	All I All I Live and All I all	
	<220>	
	<223> Description of Artificial Sequence: PepA/amdS	
	<400> 7	
15	ttcccaggat tgaggcattt tgaccacgag att	33
		-
	<210> 8	
	<211> 17	
20	<212> DNA	
	<213> Artificial Sequence	
	<220>	
	<pre><223> Description of Artificial Sequence: MBL1213</pre>	
25	· · · · · · · · · · · · · · · · · · ·	
	<400> 8	
	taacttccac cgaggtc	17
30	<210> 9	
	<211> 18 <212> DNA	
	<212> DNA <213> Artificial Sequence	
35	<220>	
	<223> Description of Artificial Sequence: Primer 122958	
	<400> 9	
	cgatcgatga ctgcctgt	18
40		
	<210> 10	
	<211> 18	
	<212> DNA	
45	<213> Artificial Sequence	
	<220>	
	<223> Description of Artificial Sequence: Primer 122956	
50	400. 10	
50	<400> 10 agagacacat agtgcctt	18
		10
	.010 17	
55	<210> 11 <211> 18	
	<211> 10 <212> DNA	
	<213> Artificial Sequence	
	<220>	
50	<pre><220> <223> Description of Artificial Sequence: Primer 122959</pre>	

```
<400> 11
     gcttatagtc gatagcgc
                                                                         18
     <210> 12
     <211> 18
     <212> DNA
     <213> Artificial Sequence
10
     <220>
     <223> Description of Artificial Sequence: Primer 122960
     <400> 12
     cctctccca gcgatggt
                                                                         18
15
     <210> 13
     <211> 18
     <212> DNA
20
     <213> Artificial Sequence
     <220>
     <223> Description of Artificial Sequence: Primer122962
25
     <400> 13
     atggaataca tactgctt
                                                                         18
     <210> 14
30
     <211> 18
     <212> DNA
     <213> Artificial Sequence
35
     <223> Description of Artificial Sequence: Primer122961
     <400> 14
     atgaaaccca ctgtagct
                                                                         18
40
     <210> 15
     <211> 18
     <212> DNA
     <213> Artificial Sequence
45
     <223> Description of Artificial Sequence: Primer 122963
     <400> 15
     tgctcgataa gcgggtcc
50
                                                                         18
     <210> 16
     <211> 18
55
     <212> DNA
     <213> Artificial Sequence
     <223> Description of Artificial Sequence: Primer 122964
60
     <400> 16
```

	attettatgg accegett	18
5	<210> 17 <211> 20 <212> DNA <213> Artificial Sequence	
10	<pre><220> <223> Description of Artificial Sequence: Primer 124289</pre>	
	<400> 17 ccccgggaaa caagaacagg	20
15		
13	<210> 18 <211> 20 <212> DNA	
	<213> Artificial Sequence	
20	<220>	
	<223> Description of Artificial Sequence: Primer 124290	
25	<400> 18 gttggcggac cttgactatg	20
	<210> 19	
30	<211> 22 <212> DNA	
50	<213> Artificial Sequence	
	<220>	
	<pre><223> Description of Artificial Sequence: Primer 125112</pre>	
35		
	<400> 19 acagctacag tgggtttcat ct	22
		22
40	<210> 20	
	<211> 19	
	<212> DNA	
	<213> Artificial Sequence	
45	<220> <223> Description of Artificial Sequence:Primer 125111	
	<400> 20 .	
= 0	agtcaacggg ggaagtctc	19
50		
	<210> 21	
	<211> 20 <212> DNA	
55	<213> Artificial Sequence	
	<220>	
	<pre><220> <223> Description of Artificial Sequence: Primer 128330</pre>	
	•	
50	<400> 21 ctagcagcgt atcggtcagc	20

```
<210> 22
     <211> 19
     <212> DNA
     <213> Artificial Sequence
     <220>
     <223> Description of Artificial Sequence: Primer 130887
10
     <400> 22
     cttggaaaag aaacgatag
                                                                         19
15
     <210> 23
     <211> 19
     <212> DNA
     <213> Artificial Sequence
20
     <220>
     <223> Description of Artificial Sequence: Primer 130888
     <400> 23
     aacgtacgct ttcctcctt
                                                                         19
25
     <210> 24
     <211> 21
     <212> DNA
30
     <213> Artificial Sequence
     <223> Description of Artificial Sequence: Primer 134135
35
     <400> 24
     gggtccgtcc agtccgttct t
                                                                         21
     <210> 25
40
     <211> 24
     <212> DNA
     <213> Artificial Sequence
     <220>
45
     <223> Description of Artificial Sequence: Primer -48
          reverse
     <400> 25
     agcggataac aatttcacac agga
                                                                         24
50
     <210> 26 ·
     <211> 17
     <212> DNA
55
     <213> Artificial Sequence
     <220>
     <223> Description of Artificial Sequence: Primer-40
           Universal
60
     <400> 26
```

	gttttcccag tcacgac	17
5	<210> 27 <211> 19 <212> DNA <213> Artificial Sequence	
10	<220> <223> Description of Artificial Sequence: Primer PstI	
	<400> 27 teatecetgg tgttactge	19
15		
	<210> 28 <211> 17	
	<212> DNA	
20	<213> Artificial Sequence	
20	<220>	
	<223> Description of Artificial Sequence: PstII	
	<400> 28	
25	catggattgg ctggccg	17
	<210> 29	
	<211> 18	
30	<212> DNA <213> Artificial Sequence	
	12107 ALCITICIAL Bequence	
	<220>	
35	<223> Description of Artificial Sequence: Prt270n	
	<400> 29	
	tactctccag attgcctg	18
40	<210> 30	
	<211> 18 <212> DNA	
	<213> Artificial Sequence	
45	<220>	
1.5	<223> Description of Artificial Sequence: Prt1420r	
	-	
	<400> 30 tgagatacca ctcagcag	18
50	5.5	10
	<210> 31	
	<211> 18	
c c	<212> DNA	
55	<213> Artificial Sequence	
	<220>	
	<223> Description of Artificial Sequence: Prt1350n	
60	<400> 31	
	tgcacttctc tgtctctg	18

```
<210> 32
     <211> 18
     <212> DNA
     <213> Artificial Sequence
     <223> Description of Artificial Sequence: Prt2365r
10
     <400> 32
     gacttctggc atcagttg
                                                                         18
15
     <210> 33
     <211> 18
     <212> DNA
     <213> Artificial Sequence
20
     <220>
     <223> Description of Artificial Sequence: Prt2320n
     ctcatggatg gcatgatc
                                                                         18
25
     <210> 34
     <211> 33
     <212> PRT
30
     <213> Aspergillus niger
     <220>
     <221> ZN_FING
     <222> (1)..(33)
35
     <400> 34
     Met Thr Ala Cys His Thr Cys Arg Lys Leu Lys Thr Arg Cys Asp Leu
40
     Asp Pro Arg Gly His Ala Cys Arg Arg Cys Leu Ser Leu Arg Ile Asp
                  20
                                       25
     Cys
45
     <210> 35
     <211> 20
50
     <212> DNA
     <213> Artificial Sequence
     <220>
     <223> Description of Artificial Sequence: Degenerated
55
           Primer 137396
     <400> 35
     atgacygcyt gycayacytg
                                                                         20
60
     <210> 36
```

```
<211> 20
     <212> DNA
     <213> Artificial Sequence
     <220>
     <223> Description of Artificial Sequence: Degenerated
           Primer 137397
     <400> 36
10
     arrcancgnc greargertg
                                                                        20
     <210> 37
     <211> 20
     <212> DNA
     <213> Artificial Sequence
     <223> Description of Artificial Sequence: Primer 137394
20
     <400> 37
     atgactgcct gtcacacatg
                                                                        20
25
     <210> 38
     <211> 20
     <212> DNA
     <213> Artificial Sequence
30
     <223> Description of Artificial Sequence: Primer 137395
     agacagcgac ggcacgcatg
                                                                        20
35
     <210> 39
     <211> 21
     <212> DNA
40
    <213> Artificial Sequence
     <223> Description of Artificial Sequence: Primer 144428
     <400> 39
     caccgagttt taagcttgcg g
                                                                        21
     <210> 40
50
     <211> 20
     <212> DNA
     <213> Artificial Sequence
55
    <223> Description of Artificial Sequence: Primer 144429
     <400> 40
    gcgatcttga tccacgaggg
                                                                       20
б0
     <210> 41
```

```
<211> 21
      <212> DNA
      <213> Artificial Sequence
  5
      <220>
      <223> Description of Artificial Sequence: Primer 153468
      cgggatgaat tgtagagagg c
                                                                          21
 10
      <210> 42
      <211> 25
      <212> DNA
 15
      <213> Artificial Sequence
      <223> Description of Artificial Sequence: Primer 140358
. 20
      <400> 42
      cgcaagctta aaactcggtg cgatc
                                                                          25
      <210> 43
 25
      <211> 20
      <212> DNA
      <213> Artificial Sequence
      <220>
 30
      <223> Description of Artificial Sequence: Primer 140359
      <400> 43
      cctcgtggat caagatcgca
                                                                          20
 35
      <210> 44
      <211> 22
      <212> DNA
      <213> Artificial Sequence
 40
      <220>
      <223> Description of Artificial Sequence: Primer 175653
      <400> 44
 45
      gatgaaaaga ataatcggcg ag
                                                                          22
      <210> 45
      <211> 21
      <212> DNA
 50
      <213> Artificial Sequence
      <223> Description of Artificial Sequence: Primer 175654
 55
      <400> 45
      cgcggcacac taccccgtt g
                                                                          21
 60
      <210> 46
      <211> 22
```

```
<212> DNA
     <213> Artificial Sequence
     <220>
    <223> Description of Artificial Sequence: Primer B043G08
     <400> 46
    atctagetea ageattageg ge
                                                                     22
10
     <210> 47
     <211> 21
     <212> DNA
     <213> Artificial Sequence
15
     <223> Description of Artificial Sequence: Primer
          B0403G09
20
    <400> 47
    aatttcggcc ctttagtgtc c
                                                                     21
    <210> 48
25
    <211> 2931
     <212> DNA
    <213> Aspergillus oryzae
    <220>
30
    <221> intron
    <222> (1028)..(1135)
     <220>
     <221> intron
35
    <222> (1538)..(1591)
    <220>
    <221> intron
    <222> (2018)..(2066)
40
    <220>
    <221> intron
    <222> (2297)..(2347)
    <400> 48
    gatateteat gatetgegtg ateggettge etectatett agateaeceg ggettettea 60
    aatcagcaac aacgctcaga catgtcccct gagaggtgat ccaaatcata cacgagagaa 120
    cgcggaaacg caaattaagg atgagcgaaa aagagaaaaa aatccgttgt tcctgagtca 180
    tgacgaatga gcaaaagtca aacacactt ctgcttttgg ggggtatgcc cgatcacaat 240
50
    cttcaacccg ccatgataag agacacacgc tatcgacaaa tcaccggagg tcaagattag 300
    catggaagcg ggactcccta tggagccggc ttacatcggg cgcactgcaa tggcgcacgt 420
    caatcaaccc ctctcttgtt gcagtgccta gtatgccaaa ccaccctttc tattcttcta 480
    gaaaccacac cctagagact cggatctaca cggattggtt ggaatgctcc gattagttgg 540
    catttacccc aggtcaaaat ggataatcaa tctaacggag tctatttcgt caactgcctg 600
    ccagetagea caateteete tteaegeeeg geegtggget gttaaaaggg teaatteeet 660
    ccccacctgt gtggattctc tatgatttgc acgggatctg acttggtttc cacaattctt 720
    cttgctctca gcttgttcta ctcgccgatt attcttttca tcaacgcggc acactacccc 780
    cgttgtctga tgtcatgact agaactactg ttgaacctat caaatatgag gccccttcgt 840
    gggagcataa gagcgtgcat gtgtccgacg accacaggag aatcatcccc aatgtcggcg 900
60
    acgacgcgac gcgcccaaag ggccgcatta gacgttcaat gaccgcttgt aatacctgcc 960
```

```
gcaagettaa aacteggtge gatettgate caegagggea tgcatgeegg eggtgtetat 1020
     ctttaaggtc gggtgccacc gttatccact ttgtcaaatc tcttacgtca aaatggggga 1080
     tcccatgtcc tgccaagacc aaataagcct ttcttgagta ctaatgtttc tataggatcg 1140
     actyteaget eccegagacg agtgageget tteaggacag tactecaatg tggteagacg 1200
     caacgacagc tateccetec ategaggage gteteaette cetagagagg agtatgagag 1260
     agatgaccgg catgettegg cagatettga atcaatcacc aagegtetet aatateteeg 1320
     teceteeget ageteggagt gtteataegg aagaaaegge etecattgaa ggaaaeteat 1380
     teggteettt cetacetaaa eeegttegge taatteagga eeteeaatet gagttttttg 1440
     gggagacaaa ccgcatccct gttgaatctc ctttcttggg taacagtttt gagaagggta 1500
10
     tettagatte taagttgtet eteaagttgg tacagetgta tggteacteg teatgteeat 1560
     ctgcctctat agccgctaat gcttgagcta gatttgtgga taatttcggc cctttagtgt 1620
     ccataaataa tcagtcggac ttccacaacg agatgaggaa caccgattcg ttgttatata 1680
     gtactgcctg tcttctggcc tcccgatatg tgccaggcat accaccaccg attgtccata 1740
     ccatgaacct ccaagttcga cataaggcag tcaatctgct gtgggaagaa ccgcctttga 1800
15
     aatacgaatc gctccaggca ctcgcccttc tttgtttatg gccagcggcg ggtcaaaagg 1860
     agttccccat agatggctgg ttactgagcg ggactgcaat caatcatgcc ctcgtctcct 1920
     ttgacttcct caatcatgtg ccttcagagc ttctcattga taacgatatc gccgctcaat 1980
     tgcggctctg gaacgctttc tgtttaacac agttacagta ggtacaacat ttccggctta 2040
     actccaactt gctaatgcag aaatagtttc gctgttggca acgcacgtcc attccattta 2100
20
     ccacagagat atctcgatta ttgcccacga cttcttgagc accccgctgc aacagttgag 2160
     gatggcaagg tcgtagcaga gatccagttg tacttgatca cattgcgact ccaagccaac 2220
     gagcaacgta tgcgattcgc ggaggttgaa tacgaagaga ttgaacgatg gaaagttgaa 2280
     tgggcccatc ttcttggtaa ggttaagcaa cgaggaccat ctcatataaa tgctaactat 2340
     tcaacagetg gtgatgaaaa ttcaacattt gagettagte tetggttetg tcaaateete 2400
     ctgcatcgga cagcaatgag gttccaagcg gagtctgaga gactcacgtc ggaaattctc 2460
     caaggatege gettgateat etegaaatte etgeaactee gatttgteac egetetaaga 2520
     gtggtcgatc aggcgtactt catcgtcggt tatgccgctc taaatctttg cgacttcaac 2580
     ttcctcgacc ccctcattga ccagatccag atgtttctgc tgcatctgtc gccaaacgaa 2640
     gaccacateg cataceggtt ttegtgeatg atageegagt teaagegteg etgtgeegaa 2700
30
     tgcaacgacc cttgcagcgc agtcgacggt tctcaatgct cgttcggaga tgcccggaag 2760
     atgagcatgg aacaggtaca attcgtgcca ccactagtag atagcatgat tgggggatat 2820
     agcgctctgg aacagctgat ccctgaggtc atgccacact catttccgga aagtgtcata 2880
     agtggcatgg ctgtgactga agccatccct gtgggatcgg cgccatacta g
35
     <210> 49
     <211> 624
     <212> PRT
     <213> Aspergillus oryzae
40
     <400> 49
     Met Thr Arg Thr Thr Val Glu Pro Ile Lys Tyr Glu Ala Pro Ser Trp
45
     Glu His Lys Ser Val His Val Ser Asp Asp His Gly Arg Ile Ile Pro
     Asn Val Gly Asp Asp Ala Thr Arg Pro Lys Gly Arg Ile Arg Arg Ser
50
     Met Thr Ala Cys Asn Thr Cys Arg Lys Leu Lys Thr Arg Cys Asp Leu
     Asp Pro Arg Gly His Ala Cys Arg Arg Cys Leu Ser Leu Arg Ile Asp
55
     Cys Gln Leu Pro Glu Thr Ser Glu Arg Phe Gln Asp Ser Thr Pro Met
60
     Trp Ser Asp Ala Thr Thr Ala Ile Pro Ser Ile Glu Glu Arg Leu Thr
                                                         110
```

	Ser	Leu	Glu 115	Arg	Ser	Met	Arg	Glu 120	Met	Thr	Gly	Met	Leu 125	Arg	Gln	Ile
5	Leu	Asn 130	Gln	Ser	Pro	Ser	Val 135	Ser	Asn	Ile	Ser	Val 140	Pro	Pro	Leu	Ala
10	Arg 145	Ser	Val	His	Thr	Glu 150	Glu	Thr	Ala	Ser	Ile 155	Glu	Gly	Asn	Ser	Phe 160
10	Gly	Pro	Phe	Leu	Pro 165	Lys	Pro	Val	Arg	Leu 170	Ile	Gln	qaA	Leu	Gln 175	Ser
15	Glu	Phe	Phe	Gly 180	Glu	Thr	Asn	Arg	Ile 185	Pro	Val	Glu	Ser	Pro 190	Phe	Leu
	Gly	Asn	Ser 195	Phe	Glu	Lys	Gly	Ile 200	Leu	Asp	Ser	Lys	Leu 205	Ser	Leu	Lys
20	Leu	Val 210	Gln	Leu	Phe	Val	Asp 215	Asn	Phe	Gly	Pro	Leu 220	Val	Ser	Ile	Asn
25	Asn 225	Gln	Ser	Asp	Phe	His 230	Asn	Glu	Met	Arg	Asn 235	Thr	Asp	Ser	Leu	Leu 240
25	Tyr	Ser	Thr	Ala	Сув 245	Leu	Leu	Ala	Ser	Arg 250	Tyr	۷al	Pro	Gly	Ile 255	Pro
30	Pro	Pro	Ile	Val 260	His	Thr	Met	Asn	Leu 265	Gln	Val	Arg	His	Lув 270	Ala	Val
	Asn	Leu	Leu 275	Trp	Glu	Glu	Pro	Pro 280	Leu	Lys	Tyr	Glu	Ser 285	Leu	Gln	Ala
35	Leu	Ala 290	Leu	Leu	Cys	Leu	Trp 295	Pro	Ala	Ala	Gly	Gln 300	Lys	Glu	Phe	Pro
40	Ile 305	qsA	Gly	Trp	Leu	Leu 310	Ser	Gly	Thr	Ala	Ile 315	Asn	His	Ala	Leu	Val 320
10	Ser	Phe	Asp	Phe	Leu 325	Asn	His	Val	Pro	Ser 330	Glu	Leu	Leu	Ile	Asp 335	Asn
45	qzA	Ile	Ala	Ala 340	Gln	Leu	Arg	Leu	Trp 345	Asn	Ala	Phe	Сув	Leu 350	Thr	Gln
	Leu	His	Phe 355	Ala	Val	Gly	Asn	Ala 360	Arg	Pro	Phe	His	Leu 365	Pro	Gln	Arg
50	Tyr	Leu 370	qaA	Tyr	Сув	Pro	Arg 375	Leu	Leu	Glu	His	Pro 380	Ala	Ala	Thr	Val
55	Glu 385	Asp	Gly	Lys	Val	Val 390	Ala	Glu	Ile	Gln	Leu 395	Tyr	Leu	Ile	Thr	Leu 400
	Arg	Leu	Gln	Ala	Asn 405	Glu	Gln	Arg	Met	Arg 410	Phe	Ala	Glu	Val	Glu 415	Tyr
60	Glu	Glu	Ile	Glu 420	Arg	Trp	Lys	Val	Glu 425	Trp	Ala	His	Leu	Leu 430	Ala	Gly

aggtacagaa gtccaattgc ttccgatctg gtaaaagatt cacgagatag taccttctcc 720 gaagtaggta gagcgagtac ccggcgcgta agctccctaa ttggcccatc cqqcatctgt 780 agggcgtcca aatatcgtgc ctctcctgct ttgcccggtg tatgaaaccg gaaaggccgc 840 tcaggagetg gccageggcg cagaceggga acacaagetg gcagtegace cateeggtge 900 tctgcactcg acctgctgag gtccctcagt ccctggtagg cagctttgcc ccgtctgtcc 960 gcccggtgtg tcggcggggt tgacaaggtc gttgcgtcag tccaacattt gttgccatat 1020 ttteetgete teeceaceag etgtagatet tggtggegtg aaacteeege acetettegg 1080 ccagegeett gtagaagege gtatggette gtaceeegge cateaacaeg egtetgegtt 1140 cgaccagget gegegttete geggecatag caaccgacgt acggegttge geeetegeeg 1200 gcagcaagaa gccacggaag teegeeegga gcagaaaatg cccacgctac tgcgggttta 1260 tatagacggt ccccacggga tggggaaaac caccaccacg caactgctgg tggccctggg 1320 ttegegegae gatategtet aegtaeeega geegatgaet taetggeggg tgetggggge 1380 ttccgagaca atcgcgaaca tctacaccac acaacaccgc ctcgaccagg gtgagatatc 1440 ggccggggac gcggcggtgg taatgacaag cgcccagata acaatgggca tgccttatgc 1500 15 cgtgaccgac gccgttctgg ctcctcatat cgggggggag gctgggagct cacatgcccc 1560 geocceggee eteacectea tettegaceg coateceate geogeoctec tgtgctaccc 1620 ggccgcgcgg taccttatgg gcagcatgac cccccaggcc gtgctggcgt tcgtggccct 1680 catecegecg acettgeecg geaceaacat egtgettggg geeetteegg aggacagaca 1740 categacege etggecaaac gecagegeec eggegagegg etggacetgg etatgetgge 1800 20 tgcgattcgc cgcgtttacg ggctacttgc caatacggtg cggtatctgc agtgcggcgg 1860 gtcgtggcgg gaggactggg gacagettte ggggaeggee gtgeegeece agggtgeega 1920 gccccagagc aacgcgggcc cacgacccca tatcggggac acgttattta ccctgtttcg 1980 gggccccgag ttgctggccc ccaacggcga cctgtataac gtgtttgcct gggccttgga 2040 cgtcttggcc aaacgcctcc gttccatgca cgtctttatc ctggattacg accaatcgcc 2100 cgccggctgc cgggacgccc tgctgcaact tacctccggg atggtccaga cccacgtcac 2160 caccccegge tecatacega egatatgega eetggegege aegtttgeee gggagatggg 2220 ggaggctaac tgaaacacgg aaggagacaa taccggaagg aacccgcgct atccggatcc 2280 acttaacgtt actgaaatca tcaaacagct tgacgaatct ggatataaga tcgttggtgt 2340 cgatgtcagc tccggagttg agacaaatgg tgttcaggat ctcgataaga tacgttcatt 2400 30 tgtccaagca gcaaagagtg ccttctagtg atttaatagc tccatgtcaa caagaataaa 2460 acgcgttttc gggtttacct cttccagata cagctcatct gcaatgcatt aatgcattga 2520 ctgcaaccta gtaacgcctt caggctccgg cgaagagaag aatagcttag cagagctatt 2580 ttcattttcg ggagacgaga tcaagcagat caacggtcgt caagagacct acgagactga 2640 ggaatccgct cttggctcca cgcgactata tatttgtctc taattgtact ttgacatgct 2700 35 cctcttcttt actctgatag cttgactatg aaaattccgt caccagccct gggttcgcaa 2760 agataattgc atgtttcttc cttgaactct caagcctaca ggacacacat tcatcgtagg 2820 tataaacctc gaaatcattc ctactaagat ggtatacaat agtaaccatg catggttgcc 2880 tagtgaatgc tccgtaacac ccaatacgcc ggccgaaact tttttacaac tctcctatga 2940 gtcgtttacc cagaatgcac aggtacactt gtttagaggt aatccttctt tctagaagtc 3000 40 ctcgtgtact gtgtaagege ccactecaca tetecacteg acetgeagge atgcaagett 3060 ggcgtaatca tggtcatagc tgtttcctgt gtgaaattgt tatccgctca caattccaca 3120 caacatacga gccggaagca taaagtgtaa agcctggggt gcctaatgag tgagctaact 3180 cacattaatt gcgttgcgct cactgcccgc tttccagtcg ggaaacctgt cgtgccagag 3240 eggeegetet geattaatga ateggeeaac gegeggggag aggeggtttg egtattggge 3300 45 getetteege tteetegete aetgaetege tgegeteggt egtteggetg eggegagegg 3360 tatcagetca etcaaaggeg gtaataeggt tatceacaga atcaggggat aaegeaggaa 3420 agaacatgtg agcaaaaggc cagcaaaagg ccaggaaccg taaaaaggcc gcgttgctgg 3480 cgtttttcca taggctccgc ccccctgacg agcatcacaa aaatcgacgc tcaagtcaga 3540 ggtggcgaaa cccgacagga ctataaagat accaggcgtt tccccctgga agctccctcg 3600 tgcgctctcc tgttccgacc ctgccgctta ccggatacct gtccgccttt ctcccttcgg 3660 gaagcgtggc gctttctcat agctcacgct gtaggtatct cagttcggtg taggtcgttc 3720 getecaaget gggetgtgtg cacgaaceee cegtteagee egacegetge geettateeg 3780 gtaactateg tettgagtee aacceggtaa gacacgaett ategecaetg geageageea 3840 ctggtaacag gattagcaga gcgaggtatg taggcggtgc tacagagttc ttgaagtggt 3900 55 ggcctaacta cggctacact agaaggacag tatttggtat ctgcgctctg ctgaagccag 3960 ttaccttcgg aaaaagagtt ggtagctctt gatccggcaa acaaaccacc gctggtagcg 4020 gtggtttttt tgtttgcaag cagcagatta cgcgcagaaa aaaaggatct caagaagatc 4080 ctttgatett ttetaegggg tetgaegete agtggaacga aaacteaegt taagggattt 4140 tggtcatgag attatcaaaa aggatcttca cctagatcct tttaaattaa aaatgaagtt 4200 ttaaatcaat ctaaagtata tatgagtaaa cttggtctga cagttaccaa tgcttaatca 4260 gtgaggcacc tatctcagcg atctgtctat ttcgttcatc catagttgcc tgactccccg 4320

```
tegtgtagat aactaegata egggagget taccatetgg eeccagtget geaatgatac 4380
     cgcgagaccc acgctcaccg gctccagatt tatcagcaat aaaccagcca gccggaaggg 4440
     ccgagcgcag aagtggtcct gcaactttat ccgcctccat ccagtctatt aattgttqcc 4500
     gggaagctag agtaagtagt togcoagtta atagtttgcg caacgttgtt gccattgcta 4560
     caggeategt ggtgteaege tegtegtttg gtatggette atteagetee ggtteeeaac 4620 gateaaggeg agttacatga teececatgt tgtgcaaaaa ageggttage teetteggte 4680
     ctccgatcgt tgtcagaagt aagttggccg cagtgttatc actcatggtt atggcagcac 4740
     tgcataattc tcttactgtc atgccatccg taagatgctt ttctgtgact ggtgagtact 4800
     caaccaagtc attctgagaa tagtgtatgc ggcgaccgag ttgctcttgc ccggcgtcaa 4860
10
     tacgggataa taccgcgcca catagcagaa ctttaaaagt gctcatcatt ggaaaacgtt 4920
     ctteggggcg aaaactctca aggatcttac cgctgttgag atccagttcg atgtaaccca 4980
     ctcgtgcacc caactgatct tcagcatctt ttactttcac cagcgtttct gggtgagcaa 5040
     aaacaggaag gcaaaatgcc gcaaaaaagg gaataagggc gacacggaaa tgttgaatac 5100
     tcatactctt cctttttcaa tattattgaa gcatttatca gggttattgt ctcatgagcg 5160
     gatacatatt tgaatgtatt tagaaaaata aacaaatagg ggttccgcgc acatttcccc 5220
     gaaaagtgcc acctgacgtc taagaaacca ttattatcat gacattaacc tataaaaaata 5280
     ggcgtateac gaggcccttt cgtctcgcgc gtttcggtga tgacggtgaa aacctctgac 5340
     acatgcaget eceggagacg gtcacagett gtctgtaage ggatgceggg agcagacaag 5400
     cccgtcaggg cgcgtcagcg ggtgttggcg ggtgtcgggg ctggcttaac tatgcggcat 5460
20
     cagagcagat tgtactgaga gtgcaccata tcgacgctct cccttatgcg actcctgcat 5520
     taggaagcag cccagtagta ggttgaggcc gttgagcacc gccgccgcaa ggaatggtgc 5580 atgcaaggag atggcgcca acagtccccc ggccacgggg cctgccacca tacccacgcc 5640
     gaaacaagcg ctcatgagcc cgaagtggcg agcccgatct tccccatcgg tgatgtcggc 5700
     gatataggcg ccagcaaccg cacctgtggc gccggtgatg ccggccacga tgcgtccggc 5760
25
     gtagaggate tggetagega tgaccetget gattggtteg etgaccattt ceggggtgeg 5820
     gaacggcgtt accagaaact cagaaggttc gtccaaccaa accgactctg acggcagttt 5880
     acgagagaga tgatagggtc tgcttcagta agccagatgc tacacaatta ggcttgtaca 5940
     tattgtcgtt agaacgcggc tacaattaat acataacctt atgtatcata cacatacgat 6000
     ttaggtgaca ctata
30
     <210> 51
     <211> 18
     <212> DNA
35
     <213> Artificial Sequence
     <223> Description of Artificial Sequence: Primer
           B1042E05
40
     <400> 51
     cgcgcgtatc ctattgcc
                                                                            18
     <210> 52
     <211> 20
     <212> DNA
     <213> Artificial Sequence
50
     <400> 52
     gccggaaatg ttgtacctac
                                                                            20
     <210> 53
55
     <211> 80
     <212> DNA
     <213> Artificial Sequence
     <220>
60
     <223> Description of Artificial Sequence: ICA217
```

	3,333	60 80
5		
	<210> 54	
	<211> 80	
	<212> DNA	
	<213> Artificial Sequence	
LO	· ·	
	<220>	
	<223> Description of Artificial Sequence: ICA218	
	<400> 54	
L5		60 80

INTERNATIONAL SEARCH REPORT

Inte al Application No PCT/DK 01/00169

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12N15/31 C12N15/67 //(C12N15/31,C12R1:69) According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C12N IPC 7 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ° Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. WO 00 20596 A (NOVONORDISK AS) 13 April 2000 (2000-04-13) P,X 1-50 the whole document χ DATABASE SWISS-PROT [Online] 1,5,12, accession no. P39529 14,15, 1 November 1997 (1997-11-01) 17 - 20,25B PURNELLE ET AL: "Putative 86.7 kda transcriptional regulatory protein in NUC1-NCE1 intergenic region" XP002901831 page 45 -page 76 Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the International search report 20. 08. 2001 2 August 2001 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+31-70) 340-3016 Hampus Rystedt

INTERNATIONAL SEARCH REPORT

Int Inal Application No PCT/DK 01/00169

		LC1/DK 01/00103
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with Indication, where appropriate, of the relevant passages	Relevant to claim No.
X	DATABASE SWISS-PROT [Online] accession no P38114, 1 February 1995 (1995-02-01) ENTIAN KD ET AL: "Putative 126.9 kda transcriptional regulatory protein in YSW1-RIB7 intergenic region" XP002901832 page 106 -page 142	1,5,12, 14,15, 17-20,25
A	WO 97 12045 A (CHIRON CORP) 3 April 1997 (1997-04-03) the whole document	1-50
A .	WO 95 35385 A (NOVONORDISK AS) 28 December 1995 (1995-12-28) the whole document	1-50
A	K L PETERSEN: "A new transcriptional activator for amylase genes in Aspergillus" MOL GEN GENET, vol. 262, 1999, pages 668-676, XP002901833 the whole document	1-50

INTERNATIONAL SEARCH REPORT

Information on patent family members

Int onal Application No PCT/DK 01/00169

Patent document cited in search report	Patent document cited in search report		Patent family member(s)		Publication date
WO 0020596	A	13-04-2000		850999 A 117798 A	26-04-2000 25-07-2001
WO 9712045	A	03-04-1997	AU 7	112796 A	17-04-1997
WO 9535385	Α	28-12-1995	CN 1 EP 0 FI JP 10	733895 A 150824 A 770139 A 965031 A 501414 T 025185 A	15-01-1996 28-05-1997 02-05-1997 16-12-1996 10-02-1998 15-02-2000
			~		